

FIGURE 1

ACTGCACCTCGGTTATCGATTGAATCCCCGGGATCCTCTAGAGATCCCTGACCTCGA
CCCACCGCGTCCGGGCCGGAGCAGCACGGCCGAGGACCTGGAGCTCCGGCTGCGCTTCCG
CAGCGCTACCCGCCATGCGCTGCGCGCCGGCTGGGCTCCTGCCGCTTGTGCTG
CTGCTGCCCGCGCCGGAGGCCAAGAACGCCGACGCCCTGCCACGGTGCGCTGGGCT
GGTGGACAAGTTAACCAAGGGGATGGTGGACACCGCAAAGAACAGAACTTTGGCGGCGGAAACA
CGGCTTGGGAGGAAAAGACGCTGTCCAAGTAGCAGTCCAGCGAGATTGCCCTGCTGGAGATC
CTGGAGGGGCTGTGCGAGAGCAGCAGTCGAATGCAATCAGATGCTAGAGGCGCAGGAGGA
GCACCTGGAGGCCTGGCTGCGACTGAAAGAGCGAATATCCTGACTTATCGAGTGGTTT
GTGTGAAGACACTGAAAGTGTGCTGCTCTCCAGGAACCTACGGTCCGACTGTCGCATGC
CAGGGCGGATCCCAAGAGGCCCTGCGAGGGAAATGCCACTGAGCGGAGATGGAGCAGACA
GGCGCAGGGTCTGCGGCTGCCAACATGGGTACCAAGGCCGCTGTGACTGACTGCCATGG
ACGGCTACTTCAGCTCGCTCCGAACAGAACCCACAGCATCTGCACAGCCTGTGACGAGTCC
TGCAAGACGTGCTCGGGCTGACCAACAGAGACTGCGCGAGTGTGAAGTGGCTGGGTGCT
GGACGAGGGCGCTGTGTTGGATGTGGACGAGTGTGCGGCCAGGCCCTCTGCAGCGCTG
CGCAGTTCTGTAAGAACGCAACGGCTCTACACGTGCGAAGAGTGTGACTCCAGCTGTG
GGCTGCACAGGGAAAGGCCAGGAAACTGTAAAGAGTGTATCTCTGGCTACCGAGGGACA
CGGACAGTGTGACATGTGGACGAGTGTGACTAGCAGAAAAAAACCTGTGAGGAAAAACG
AAAACGTCTACAATACTCCAGGGAGCTACGCTGTGTCCTGACGGCTCGAAGAACG
GAAGATGCCCTGTGCCGCCAGAGGCTGAAGCCACAGAAGGGAAAGCCGACACAGT
GCCCTCCCGCGAACACTGTAAGTGCCGACTTACCTTAAATTATTCAAGGATGTC
CGTGGAAATGTGCCCTGAGGATGCCGTCTCTGCAGTGGACAGCGGGGGAGAGGCTG
CTGCTCTCTAACGGTTGATTCTCATTTGCCCTAAACAGCTGCATTCTGGTTGTTCTTA
AACAGACTGTATTTGATACAGTTCTTGTAAAAATTGACCATTGAGGTAAATCAGG
AGGAAAAAAAGGGCGGCCGGACTCTAGAGTCGACCTGAGAAC
TTGCCGCCATGCCCAACTGTTATTGAGCTTAAATGGTACAAATAAGCAATAGCA
TCACAAATTTCACAAATAAACGATTTTTCACTGCATTCTAGTTGTGGTTGTCCAACACTC
ATCAATGTATCTTATCATGTCTGGATCGGAATTAAATTGCCGAGCAGCACCAGGCTGAAAT
AACCTGAAAGAGGAACTTGGTTAGGTACCTTCTGAGGGCAAAGAACAGCTGTGGAATG
TGTGTCAGTTAGGGTGTGGAAAGTCCCCAGGCTCCCCAGCAGGCGAGAAGTATGCAAGCATGC
ATCTCAATTAGTCAGCAACCCAGTTT

FIGURE 2

><subunit 1 of 1, 353 aa, 0 stop

><MW: 38192, pI: 4.53, NX(S/T): 2

MRLPRAALGLPLLLLPPAPEAKKPTPCRRCRGLVDKFNQGMVDTAKKNFGGNTAEEKTLSKYESSEIRL
LEILEGLCCESSDFECNQMLEAQEEHLAEAWLQLKSEYEDLFEWFCVKTLKVCCSPGTYGPDCLACQGSORPCSG
NGHCSDGSRQGDGSCRCHMGYQQPLCTDCMDGYFSSLRNEHTSICTACDESCKTCGLTNRDCGECEVGWVLDE
GACVDVDECAAEPPCSAAQFCKNANGSYTCEECDSSCVGCTGEGPGNCKECISGYAREHGQCADVDECSLAEKT
CVRKNENCYNTPGSYVCVPDPGEETEDACVPPAEAETEGESPTQLPSREDL

Signal peptide:

amino acids 1-24

 **N-glycosylation sites.**

amino acids 190-194 and 251-255

 **Glycosaminoglycan attachment sites.**

amino acids 149-153 and 155-159

 **cAMP- and cGMP-dependent protein kinase phosphorylation site.**

amino acids 26-30

 **Casein kinase II phosphorylation sites.**

amino acids 58-62, 66-70, 86-90, 197-201, 210-214, 255-259, 295-299, 339-343
and 349-353

 **Tyrosine kinase phosphorylation site.**

amino acids 303-310

 **N-myristoylation sites.**

amino acids 44-50, 54-60, 55-61, 81-87, 150-156, 158-164, 164-170, 252-258 and
313-319

 **Aspartic acid and asparagine hydroxylation site.**

amino acids 308-320

 **EGF-like domain cysteine pattern signature.**

amino acids 166-178

 **Leucine zipper pattern.**

amino acids 94-116

FIGURE 3

CAGGTCAACTGCACCTCGGTTCTATGATTGAATTCCCCGGGGATCCTCTAGAGATCCCTC
GACCTCGACCCACCGTCCGCCAGGCCAGGGCACGCCAGCGCTAAACGGGAACA
GCCCTGGCTGAGGGAGCTGCAGCGCAGCAGAGTATCTGACGCCAGGGCGTAGGTGCG
GCACGAGGAGTTTCCGGCACGCCAGGGAGCTGACGCCAGGGCGTAGGTGCGCTTC
CCTGGCCGGCTCTGGGACTCCCTCTGGACAGCATGGCCGGAGGGAGGCCCTC
CGGCCGCCGCCAGGGAGAGCCTGACCTATGGATGATGCTCACCAAGGAAGAGTACTCA
TAGGATTGAGAAGATATCCTGATTGTTTCAAGGGGGAAATGGCACCTTTACACATGAT
TTCAAGAAAAGCGCAACAGAGAATGCCAGCTTCTGCAATATCCATTCCATGAATTTTAC
CTGGCAAGCTGCAGGGCAGGCAAGAATATCTATGAAATTCTGCTTGCGCTCCCTGAGATA
AAGGCATCATGGCAGATCCAACCGTCAATGTCCTCTGCTGGGACAGTGCCCTACAAGGCA
TCAGTTGTCAGTGGTTCCATGTCAGGGAAACAGGATGGGTGGCAGCATTGAAAGT
GGATGTGATTGTTATGAAATTCTGAGGCAACACCATTCTCAAACACCTCAAATGCTATCT
TCTTTAAAACATGTCACAAAGCTGAGTCCCAGGGCGGTGGCAAATGGAGGCTTTGTAAT
GAAAGACGCATCTGCAGGTGTCCTGATGGGTTCCACGGACCTCACTGTGAGAAAGCCCTTG
TACCCACAGTATGAAATGGTGTGACTTCTGGTCACTGGGATCTGCACTTGCCACCTG
GATTCTATGGAGTGAACGTGACAACAAACTGTCACCCACTGCTTAAATGGAGGAC
TGTTTCTACCCCTGGAAAATGTATTGCCCCCAGGACTAGAGGGAGAGCAGTGTGAAATCAG
CAAATGCCACACCTCTGCAATGGAGGTAAATGCAATTGGTAAAGCAATGTAAAGTGT
CCAAAGGTTACAGGGAGACCTCTGTCAAAGGCTCTGCGAGGCTGGCTGGTGCACAT
GGAACCTGCCATGAACCCAACAAATGTCAGGAAGGGTGGCATGGAAGACACTGCAA
TAAAAGGTACGAAGCCAGCCTCATACATGCCCTGAGGCCAGCAGGCCAGCTCAGGAGC
ACACGCCCTCACTAAAAGGGCGAGGAGCCGGGATCCACCTGAATCCTAACATTACATCTGG
TGAATCCGACATCTGAAACAGTTTAAGTTACACCAAGTTACAGCCTTGTAAACCTTCA
TGTGTTGAATGTCAAATAATGTTCACTACACTAAGGTTACAGCCTTGTAAACCTTCA
TCATTATAATCAGTGTGAGCTGATATTACTCTCCCTTTAAGTTTCTAAGTACGTCTGAG
CATGATGGTATAGATTCTTCTGTTCACTGCTGGGACAGATTATATTATGTCATTGA
TCAGGTTAAAATTCTCAGTGTGAGTTGCCAGATAATTTCAAAATTACATGCAATTGTT
GTCTGGGGCAGGGAAACATCAGAAAGTTAAATTGGGAAAAATCGCTAAGTCACAAGAAT
TTGGATGGTGCAGTTAATGTTGAAGTTACAGCATTTCAGGATTTTATTGTCAGATATTAGAT
GTTTGTATACATTTTAAAATTGCTCTTAAATTTTAAACTCTAACATAATATTGTCATT
TTACCAATTCTCAGAGATTCACTGTTAAACAAACAGCTTACACTGTTGAGTGGCATTT
AAACAAATATAATATTCTAAACACAATGAAATAGGGAAATATAATGTTATGAACTTTGCA
TGGCTGAAGCAATATAATATTGTAACAAACAGCTTACCTAACATAACATTGTTAT
ACTGTTGTTGATGTTAAAGGTGCTGCTTGTAGTTTTGGAAAAAAAGGGGGGGGG
CGCCATGGCCCAACTTGTATTGCACTTATAATG

FIGURE 4

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></usr/seqdb2/sst/DNA/Dnaseqs.min/ss.DNA33094
><subunit 1 of 1, 379 aa, 0 stop
><MW: 41528, pI: 7.97, NX(S/T): 2
MARRSAFPAAALWLWSILLCLLALRAEAGPPQEESLYLWIDAHQARVLIGFEEDILIVSEGK
MAPFTHDFRKAQQRMPAIPVNIHSMNFTWQAAGQAEYFYEFSLRSLDKGIMADPTVNPLL
GTVPHKASVVQVGFPCLGKQDGVAAFEVDVIVMNSEGNTILQTPQNAIFFKTCQQAECPGGC
RNGGFCNERRICECPDGFHGPHECKALCTPRCMNGGLCVTPGFCICPPGFYGVNCDKANCST
TCFNGGTCFYPGKCICPPGLEGEQCEISKCPQPCRNGGKCIIGKSCKCSKGYQGDLCSPVVC
EPGCGAHGTCHEPNKCQCQEGWHGRHCNKRYEASLIHALRPAGAQLRQHTPSLKKAERRDP
PESNYIW
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Signal peptide:

amino acids 1-28

N-glycosylation site.

amino acids 88-92, 245-249

Casein kinase II phosphorylation site.

amino acids 319-323

Tyrosine kinase phosphorylation site.

amino acids 370-378

N-myristoylation sites.

amino acids 184-190, 185-191, 189-195, 315-321

ATP/GTP-binding site motif A (P-loop).

amino acids 285-293

EGF-like domain cysteine pattern signature.

amino acids 198-210, 230-242, 262-274, 294-306, 326-338

FIGURE 5

FIGURE 6

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></usr/seqdb2/sst/DNA/Dnaseqs.min/ss.DNA33223
><subunit 1 of 1, 164 aa, 1 stop
><MW: 18359, pI: 7.45, NX(S/T) : 1
MWRCPLGLLLLLPLAGHLALGAQQGRGRRELAPGLHLRGIRDAGGRYEQDLCCRGRADDC
ALPYLGAIKYCDLFCNRTVSDCCPDFWDFCCLGVPPPFPIQGCMHGGRIYPVLGTYWDNCNR
CTCQENRQWHGGSRHDQSHQPGQLWLAGWEQPQRLLGHDPG
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N-glycosylation site.

amino acids 78-82, 161-165

Casein kinase II phosphorylation site.

amino acids 80-84, 117-121, 126-130, 169-173, 205-209, 296-300,
411-415

N-myristoylation site.

amino acids 21-27, 39-45, 44-50, 104-110, 160-164, 224-230,
269-275, 378-384, 442-448

Amidation site.

amino acids 26-30, 318-322

Eukaryotic thiol (cysteine) proteases histidine active site.

amino acids 398-409

FIGURE 7

AGGCTCCTGGCCCTTTTACAGCAAGCTNTGCNATCCGATTGTTGTCTCAAATCCA
ATTCTCTGGGACACATNACGCCGTCTTCCTTNGCCCCAGAACCTGCTGTCTGTACACCCAC
CAGCAGCAGGGCTGCCCGNTGGCGTCTCGATGGTGCCTGGTGGTTCTGCGTCGCCAGG
GNTGGTGTCTGACCACTGCTACCCCTTCTCGGGCGTGAACGAGACGAGGCTGGCCCTGCGC
CCCCCTGTATGATGCACAGCCAGCCATGGTCGGGCAAGCGCCAGGCCACTGCCACTGC
CCCAACAGCTATGTTAATAACATGACATCTACCAAGGTCACTCCTGTCTACCGCCTCGGCTC
CAACGACAAGGAGATCATGAAGGGAGCTGATGGAGAATGGCCCTGTCCAAGCCCTCATGGAGG
TGCGATGAGGACTTCTTCCTATACAAGGGAGGCATCTACAGGCCACAGCCAGTGAGCCTTGGG
AGGCCAGAGAGATAACGCCGGCATGGACCCACTCAG

FIGURE 8

GCTGCTTGCCTGTTGATGGCAGGCTGGCCCTGCAGGCCAGGCACTGCCCTGCTGTACT
CCTGCAAAGCCCAGGTGAGCACGAGGA~~T~~CTGCCCTGCAGGTGGAGAACTGCACCCAGCTGGG
GAGCACTGCTGGACCGCGCAGTCAGCTGCCCTGCACAGCTGAAAGGCTG
CAGCTTGAACTGCGTGG**AT**ACTCACAGGACTACTACGTGGCAAGAAGAACATCACGTGCT
GTGACACCAGACTTGTGCAACGCCAGCGGGGGCATGCCCTGCAGCCGGCTGCCGCCATCCTT
GCCGTGCTCCCTGCACTCGGCCCTGCTCTGGGACCCGCCAGCTATAAGGCTCTGGGGG
CCCCGCTGCAGCCCCACTGGGTGTGGTGCCCCAGGCCCTGTGCCACTCCTCACAGACCTG
GCCCACTGGGAGGCTGTCCTGGTCCATGCCCTAAGCAGTCTGACCATGTATGT
CTGCACCCCTGCCCCACCCATGCCCTCATGCCCTCTCCAGGACTCCACCCGGCAGA
TCAGCTCTAGTGAACAGATCCGCTGCAGATGGCCCTCCAACCTCTGCTGCTGTTTC
CATGGCCAGCATTCTCCACCCCTAACCCCTGTGCTCAGGCACCTCTTCCCCAGGAAGCCTT
CCCTGCCACCCATCTATGACTTGAGCCAGGTCTGGTCCGTGGTCCCCGCAACCCAGCA
GGGACAGGCACTCAGGAGGGCCAGTAAGGCTGAGATGAAGTGGACTGAGTAGAACTGGA
GGACAAGAGTCGACGTGAGTTCTGGAGTCAGAGATGGGCCCTGGAGGCCTGGAGGAA
GGGCCAGGCCTCACATTGTGGGCTCCCTGAATGGCAGCCTGAGCACAGCTAGGCCCTT
AATAAACACCTGTTGGATAAGCCAAAAAA

FIGURE 9

MTHRTTWARRTSRAVTPTCATPAGPMPCSRLPPSLRCSLHSACSGDPASYRLWGAPLQPT
LGVPQASVPLLTDLAQWEPVLVPEAHPNASLTMYVCTPVPHPDPPMALSRTPTRQISSSDT
DPPADGPNPLCCCFHGPAGFSTLNPVLRHLFPQEAFPAHPIYDLSQVWSVVSPAPSRGQALRRAQ

Signal peptide:

amino acids 1-47

N-glycosylation site.

amino acids 31-35, 74-78, 84-88

Casein kinase II phosphorylation site.

amino acids 22-26, 76-80

N-myristoylation site.

amino acids 56-60

Amidation site.

amino acids 70-74

FIGURE 10

CCCACGGTCCGAACCTCTCAGCG**ATGGGAGCCGCCGCTGCTGCCAACCTCACTCTGT**
GCTTACAGCTGCTGATTCTCTGCTGTCAAACCTCAGTACGTGAGGGACCAGGGGCCATGACC
GACCAGCTGAGCAGCGGCAGATCCCGAGTACCAACTCTACAGCAGGACCAGTGGCAAGCA
CGTGCAGGTACCGGGCGTCGATCTCGCACCGCGAGGACGGCAACAAGTTGCCAAGCT
TCATAGTGGAGACGGACAGCTTGGCAGCGGGTTCGATCAAAGGGCTGAGAGTGAGAA
TACATCTGTATGAACAAGAGGGCAAGCTCATCGGAAGGCCAGCGGGAAAGGCAAAGACTG
CGTGTTCACGGAGATCGTGTGGAGAACAACTATA CGGCCCTCCAGAACGCCCGGACAGG
GCTGGTTCATGGCCTTCACGCCAGGGGCCGGCCCCCGCCAGGCTTCCCGCAGGCCAGAAC
CAGCGCAGGCCACTTCATCAAGCGCCTCTACCAAGGCCAGCTGCCCTCCCCAACCACGC
CGAGAACAGAACGAGTCGAGTTGTGGGCTCGGCCACCCGCCAGGACCAAGCGCACAC
GGCGGCCACGCCCTCACG**T**ACTCTGGAGGCAGGGGCCAGCACGCCCTGGGCCCTCCCC
CACCCCTTCCCTCTTAATCCAAGGACTGGGCTGGGTGGCGGGAGGGAGCCAGATCCCC
GAGGGAGGACCCCTAGGGCCCGAAGCATCCGAGCCCCCAGCTGGGAAGGGCAGGCCGGT
CCCCAGGGCGGCCACGTGCCCTTCCCGACGGGTGGCAGGCCCTGGAGAGGAAC
GAGTGTACCCCTGATCTCAGGCCACCAGCCTCTGCCGCCCTCCAGCCGGCTCTGAAGCC
CGCTGAAAGGTCAAGCAGTGAAGGCCCTTGAGACAACCGCTGGAGGTGGCTGTCCCTCAAAA
TCTGCTTCTCGGATCTCCCTCAGTCTGCCCTCAGGCCCTCCAGGCCCTGGCTAGACTGTA
GGAAGGGACTTTGTTGTTGTTGTTCAAGGAAAAAGAAGGGAGAGAGAGGAAAATAG
AGGGTTGTCCACTCTCACATTCCACGCCAGGCCCTGCACCCCACTCCAGCCCC
CGGAATAAAACCATTTCCCTGC

FIGURE 11

MGAARLLPNLTLCLQLLILCCQTQYVRDQGAMTDQLSRRQIREYQLYSRTSGKHVQVTGRRI
SATAEDGNKFAKLIVETDTFGSRVRIKGAESEKYICMNKRKGKLIGKPSGKSKDCVFTEIVLE
NNYTAFQNARHEGWFMAFTRQRPRQASRSRQNQREAHFIKRLYQGQLPFPNHAEKQQKQFEF
VGSAPTRRTKRTRRPQPLT

Signal peptide:

amino acids 1-22

N-glycosylation site.

amino acids 9-13, 126-130

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 60-64

Casein kinase II phosphorylation site.

amino acids 65-69

Tyrosine kinase phosphorylation site.

amino acids 39-48, 89-97

N-myristoylation site.

amino acids 69-75, 188-194

Amidation site.

amino acids 58-62

HBGF/FGF family signature.

amino acids 103-128

FIGURE 12

ACCTGGCCATCACCTGTTGCCAGTGTGGAAAAAATTCTCCCTGTTGAATTTCGCACATGGAG
GACAGCAGCAAGAGGGCAACACAGGCTGATAAGACCAAGAGACAGCAGGAGATTATTTAC
CATACGCCCTCAGGACGTTCCCTCTAGCTGGAGTTCTGGACTCAACAGAACCCCATCCAGT
CATTGATTTCGTTTATTTTTCTTTTCTTCCACCAATTGTTATTAT
TTCCGACTTCAGAAATGGCCTACAGACCAAACTGGCCAGCCATGGGCTTTTCT
GAAGCTTGCGCTTATCTTCCCTGGGCTCTACTCACAGGCTCCAAACTCCTGGCTGCC
CTAGTGTGTGCCCTGCGACAGGAACCTTGCTACTGTAATGAGCGAAGCTTGACCTCAGTG
CCTCTGGGATCCGGAGGGCGTACCTCTACCAACAAACCAAAATTAAATTAATG
TGGATTCTCGAGAACTGCACAAATGTAAGCTGGTACACGGCTACCTGTTATGCCAACC
AACTGGACGATTCCCATGAACTTCCAAGAATGTCAGAGTTCTCCATTGCAAGGAAA
AAATTTCAGGACATTTCAGGGCTGCTCTTGCCCAGCTTGAAGCTTGAAGAGCTGCACCT
GGATGACAACCTCCATATCCACAGTGGGGGAGAGCAGGGCTCCGGGAGGTTAGGCC
TCAAATTGTTGTTTGTCAAGAATCACCTGAGCAGTGTGCTGTTGGCTTCTGTGGAC
TTGCAAGAGCTGAGAGTGGATGAAATGAAATTGCTGTCATATGCCACATGGCCTTCAGAA
TCTCACGAGCTTGGAGCTTATTGCGACGGGAAACCTCTGACCAACAGGGTATTCGCG
AGGGCACCTTCAGCCATCTCACCAAGCTCAAGGAATTTCATTTGTACTTAATTCTGCTGCC
CACCCCTCCCGATCTCCAGGTACGCATCTGATCAGGCTCTATTGCAAGGACAACAGAT
AAACACACATTCTTGTACAGCCTCTCAAAACTCTGCGTAAGCTGGAACGGCTGGATATATCA
ACAACCAACTCGGGATGCTGACTCAAGGGTTTTGTACTGAGTAAATCTCCAACTGAAAGCAGCT
ACTGCTGGAAATAACCTGGTTTGTACTGAGTAAATGGGTACAGAATGGCTCAA
ATATATCCCTCATCTCTCAACGTGGGGTTTGTGCAAGGTCTGAACAAGTCCGGG
GGATGGCGTCAAGGAAATTAAATGAAATTGCTTGTCTGTCACCCACCCGGCCTG
CCTCTTCACCCCAGCCCCAAGTACAGCTTCTGGGACACTCAGCCTCCACCCCTCTCTAT
TCCAAACCCTAGCAGAAAGCTACAGCCTCCAACCTCTACACATCGAAACTTCCACGATTC
CTGACTGGGATGCGAGGAAAGTGCACCCACCTTCTGAAACGGGATCAGCTCTATC
CATTGTGAATGATACTTCAAGTCAGCTGGCTCTCTTCAACGTGATGGCATA
CAAACCTCACATGGTAAATGGGCCACAGTTAGTAGGGGCATGTTCAAGGAGGCCATAG
TCACGGGTGAGAAGCAACACCTGAGGCTGGTTAATTAGAGCCCGATCCACCTATCGGATT
TGTTAGTGCACCTGGATGCTTAAACTACCGCGGGTAGAGACACCATTTGTTAGAGGC
CACCCCATGCTCTTATCTGAACACGGCAGAACACAGCTCCAGGCTATGAGCAGCGA
CGTCCACAGCATGGCTCCCCCTTCTGCTGGGGGCTTGTATGGGGCGCGGTGATATT
GTGCTGGGTCTGCTAGCGTCTTTGTGGCATATGACACAAAAGGGGGCTACACCTC
CCAGAAGTGGAAATAACACCGGGCGGGAAAGTGAATTGCGAGGCAGGCACCAAGA
AGGACAACCTCACCTGGAGATGACAGAAACAGTTCAAGATCTCTTAAATAACGAT
CAACTCCCTAAAGGAGATTCAAGACTGCGACGCCATTACACCCAAATGGGGCATTAATTA
CACAGACTGCCATATCCCCAACACATGCGATACTGCAACAGCAGCGTGGCAGACCTGGAGC
ACTGCGCATACGTGACAGCCAGAGGCCAGCGTTATCAAGGGCGACAATTAGACTTGGAGA
CACACTCGTGTGACATAAAGACACGCAGATTACATTGATAATGTTACACAGATGCAT
TTGTGCAATTGAATACTCTGTAATTATACGGTGTACTATATAATGGGATTAAAAAAAGTG
CTATTTCTATTCAAGTTAAATTACAAACAGTTGTAAACTCTTGCTTTAAATCTT

FIGURE 13

MGLQTTKWPSPHGAFFLKSWLIIISGLYSQVSKLLACPSVCRCDRNFVYCNERSLTSVPLGIP
EGVTVLYLHNNQINNAGFPAAELHNVQSVHTVLYGNQLDEFPMNLPKNVRVLHLQENNIQTI
SRAALAQLLKLEELHLDNSISTVGVEDGAFREAIISLKLFLSKNHSSVPVGLPVDLQELR
VDENRIAVISDMAFQNLTSLERLIVDGNNLTNKGIAEGTFSHLTKLKEFSIVRNSLSHPPPDL
LPGTHLIRLYLQDNQINHIPLTAFSNLRKLERLDISNNQLRMLTQGVFDNLNSNLKQLTARNN
PWFCDCSIKWVTEWLKYIPLLSSNLRKLERLDISNNQLRMLTQGVFDNLNSNLKQLTARNN
APSTASPTTQPPTLSIPNPSRSYTPPTPTTSKLPTIPDWDRERVTTPPISERIQLSIHFVND
TSIQVSWLSSLFTVMAYKLTWVKMGHSLVGGIVQSERIVSGEKGHQHLSLVNLEPRSTYRICLVP
DAFNYRAVEDTICSEATTHASYLNNGSNTASHEQTTSHSMSGPFLLAGLIGGAVIFVVLVVL
LSVFCWHMHKKGRYTSQKWKYNRRKDDYCEAGTKKDNSILEMTETSFQIVSLNNNDQLLKG
DFRLQPIYTPNGGINYTDCHIPEPNMRYCNSSPVDEHCHT

Signal peptide:

amino acids 1-42

Transmembrane domain:

amino acids 542-561

N-glycosylation site.

amino acids 202-206, 298-302, 433-437, 521-525, 635-639, 649-653

Casein kinase II phosphorylation site.

amino acids 204-208, 407-411, 527-531, 593-597, 598-602, 651-655

Tyrosine kinase phosphorylation site.

amino acids 319-328

N-myristoylation site.

amino acids 2-8, 60-66, 149-155, 213-219, 220-226, 294-300,
522-528, 545-551, 633-639

Amidation site.

amino acids 581-585

Leucine zipper pattern.

amino acids 164-186

Phospholipase A2 aspartic acid active site.

amino acids 39-50

FIGURE 14

ACTTGAGCAAGCGCGCGCGCGAGACAGGGCAGAGGGCAGCTGGGCTCCGCTCGCCCTCCACGAGCG
ATCCCCGAGGGAGGGCGCGGCCCTCGCGAGGGAGGGCGACGAGCGAGACCCGGTGCCTGCGCCCTGCC
TCGCTTCGCTGCTGGCCCTCTGGCTCGCCCTCTGGCTGCTGCTGCTGGAGGTCCATCTGCGAGGT
GCTTTCGCTGATCTCGCGACAGATCGCTCTCTCTCGCGAGGGCAGGGAGGGTCACTGGAGGTCCATCT
CTAGGGGAGACACGCTCGAGCCACCCCGACGCCCTCTGGAGAGGTCTGTGAGAACAAAGCGGGAGACC
TGGTTTCTCATCTTGAGACATTGGCTCTGATGTCACCGCATGTCAGGAGGTCTATCGCAAGGGACA
AGTCTCCCTCAAGACCTTCAGAGGAAGCTCGAGGGTGGAGCTGCTGAGGGATGCGGATCTGTCACGG
GCACCATGACTGGGCTGGCCATCCAGATGCCCATACTCGAGATCGGCACTTCAGAAGCAGGGGGCCGGCC
GGGAATGTGCCACGGGTCAATGATGTCAGATGGGGACCTCAGGACTCTGGCGAGGTGCTGCTA
AGGCACATGGGACACGGGCATCTTAATCTTGGCATTTGGTGGAGGGTACAGCTAACACCTTGAAGTCATTG
GGAGTGAGCCCCATGGACCATGTCCTCTTGGCCATACTTGGCAGGAGATTCAGCTGACCTCCGTGTC
AGAAGAAGTGTGAGGGCATCTTCAGGAGGGTGTGGTGTGGAGGACTACTTGTGATCAACATCCCTG
GCTCATAGCTCTGAGGTCAAACAAAGGTCATATTCTCACTCGAGCATGCACTTGAGAATTCAGGATCTG
GTGCACTGGAGGACCAACAACTGTGAGCAGCTGTGTAATGTCGGGCTCTCGTGTGAGGTCTACAGTG
GCTAGCCTCTGGAGGAGGACTGAGGAGGGTGTGGTGTGGAGGACTACTTGTGCACTGGAGTGTGAAAC
ATGAGTGTGATAATGTCATGTCCTCATTTGGCAGGAGCTTGTGAGGATTTGCTTAACCCAGATGAAAAAA
CGTGCACAAGGATCACACTGTCACACCAAACTGGCAGGAGCTTGTGAGGAGGACTGAGGAGGAGGCT
ACTACTCGGCTGCCAGGTGCTACACTCTGGACCCAACTGGCAAAACTGCGAGCCAGTGGACACTGTGAC
AGGAGGACATGGCTGTGAGCAGCTGTGTCGAACACGGGATTCCTGCTGCGAGTGTCTCAGAAGGCTCC
TCATCAAGGAGGACTCAAGACCTTCAGGCTCCGGGTGATTACTCTGGCTGAGTGCACCATGTTGTGAATACTCT
GTGTCACATGGCAGAGCATCTTGGCTCTGAGGGAGCTGGGCTCGAGGAGCTGGCAGCAGGAGCTGTG
CAAATTGGACTCTGTGAGGGGACCGGGTGTGAGGACTTGTGAGGAGGACTCTGGCTGAGGAGCTGGTGT
GCCAGTCTTGTGAGGTTATATACTCTGGAGAATGGGAACTCTGGCTGAGGAGGACTCTGGCTGAGGAGCT
ACCATGGCTGTGAAACACATTGTGAGGACTGCACTACACCTGGCTGAGGAGGATTTCGGCTCG
CTGAGGATGGGGAAACGGCTGCCAGGAGATCTGCAAAATCACCCCAATGGCTGCAACACATTTGTGTTA
ATAATTGGGAACTCTCATCTGCAAAATGTCAGAGGGATTGTCAGTGTGAGGAGCTGGCTGCAAGAAAT
GCACATGGGGCCAAATGGGGCTCTGGATCTGGAGGACTCTGGCTGAGGAGGAAATTTTGAGGTG
TGAGGAGCTTGTGACTGGAAATTAAGATCTCTGGCAAACTTCCCCAAAGCGCTCGAGTGGGGCTGCTCG
ATTCCACACAGGTCCACACAGAGTCTGGAGAATCTCACTGGACCCAAAGACATGAAAAGCCGTGGCC
ACATGAAATACATGGGAAGGGCTTGTGACTGGCTGGCTGAGAACACATGAAAAGCCGTGGCC
GAGAAGGGGGCCAGGCCCTTTCACAAAGGTGCCCCAGGCAGCATTGTGTTACCGAGGAGCGGGCTCAGGATG
ACGTCTGGAGTGGGGAGGCTTGTGAGGAGGAAATGGTGTGAGGAGGAAAGCTTACCGGATTA
AGGGGAACTACAAAGGATTGCTCTGAGGCCAAACAAAGCATCTCTTCTGAGGAGACTTCAACATGG
ATGAGATAAGTGAACAAACTCAAGAAAAGCATTGTGAGGACTCTGGAGAAGCTCCGAGTGGAGAAGCAGGACTCTCC
CAAGGGAACTGCCCCAAACCGTCCAAACGCCAAAGAACTGAGGAGGAAACACGATCATGCAAAATGIAAAACCTTATAATGT
CCCTGTTCTAAATTGGAGGTGGCAACACAGATATCTGTTGAAGAGAACATCTTACGGTCTACACAAAGCTT
CCCATTCACAAACCTTCTGGAGGCCCTTGTGAGGAGGAAACACGATCATGCAAAATGIAAAACCTTATAATGT
TCCAGAACCTTGTGCAACAGGAAGTAGAAAATTACACAGGGCTTAGAGAGAAATGACACAGGAATGGAGGCC
ACGTCTGGAGTGGGGAGGCTTGTGAGGAGGAAATGGTGTGAGGAGGAAAGCTTACCGGATTA
TGGAAAATGCTCTGGAGATACAGATGAGATTAGAAGGAAATCCGACACATTGTGACTCATTTGATACGGGATTACAA
GAACGGAGTGCAGGGCCCAAAGCTCGAGCTATTGTTAAATCAATAATGTTGAGTAAACAACTGACTG
GAACCTGTTTGTGCCACAGAGAACAGAACAGAACAGAGTATACACTAACCTGTGATAATATACCTGTGAC
TCAGAATTCTCAAGATGAAATTACCGAGGTGAGAATGAATAGCTTGTGAGGAGGAAACTCTGGACTTAA
AACTGTCTCTGGCTCATCTGGCTTAGTGTGCAATCTCATTTGACTATACGATAAAGTTGTGACAGTCTACTT
CTGAGAACACTGGCCATAGGGAAATGCTGTTTTTGTACTGGACTTAACTTGTGATAATGTTATAGGATGATAG
CATAAAATCATAGGACATAATGACTTGTGAAACAGTTGATTTTTATCAATATTAACAAATTAAACACTTCAG

FIGURE 15

MEKMLAGCFLLILGQIVLLPAEARERSRGRSISRGRRHARTHQPQTALLESSCENKRADLVFII
DSSRSVNTHDYAKVKEFIVDILQFLDIGPDVTRVGLLQVGSTVKNEFSLKTFKRSEVERAV
KRMRLSTGTMGLAIQYALNIAFSEAE GARPLR ENVPRVIMIVTDGRPQDSVAEVAAKARD
TGILIFAIVGVGQDFNTLKSIGSEPHEDHVFLVANFSIETLTSVFOKKLCTAHMCSTLEHN
CAHFCINIPGSYVCRCKQGYIILNSDQTTCTRIQDLCAMEDHNCQLCVNVPGSFVCQCYSGYA
LAEDGKRCVAVDYCASENHGCEHECVNADGSYLCQCHEGFALNPDEKTCRINYCALNKGPC
EHECVNMEESYYCRCHRGYTLDPNGKTCRVDHCAQQDHGCEQLCLNTEDSFVCQCSEGFLI
NEDLTKTCSRVDYCLLSDHGCEYCVNMDRSFACQCPEGHVLRSDGKTCALKDSCALGDHGCE
HSCVSSEDSFVCQCFEGYILREDGKTCRRKDVCAIDHGCEHICVNSDDSYTCBLEGFRILA
EDGKRCRDKDVCKSTHHGCEHICVNNNGNSYICKCSEGTVLAEDGRRCKKTEGPIDLVFFID
GSKSLGEENFEVVKQFVTGIIDS LTISPKAARVGLLQYSTQVHTEF TLRN FNSAKDMKAVA
HMKYMGKGSMGTGALKHM FERSFTQGEGARPLSTRVPR AIAVFTDGRAQDDVSEWASKAKAN
GITMYAVVGKAIEELQEIA SEPTNKHLYAEDFSTMDEISEKLKKGICCEALEDSDGRQDS
PAGE LPKTVQQPTESEP VTINI QDLLSCNSFAVQHRYLFEE DNNLLRSTQKLSHSTKPGSPL
EEKHDQCKCENLIMFQNL ANEE VRKLTQ RL EMTQ RMEALE NRL RYR

Signal peptide:

amino acids 1-23

N-glycosylation site.

amino acids 221-225

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 115-119, 606-610, 892-896

Casein kinase II phosphorylation site.

amino acids 49-53, 118-122, 149-153, 176-180, 223-227, 243-247,
401-405, 442-446, 501-505, 624-628, 673-677, 706-710, 780-784,
781-785, 819-823, 866-870

N-myristoylation site.

amino acids 133-139, 258-264, 299-305, 340-346, 453-459, 494-500,
639-645, 690-696, 752-758, 792-798

Amidation site.

amino acids 314-318, 560-564, 601-605

Aspartic acid and asparagine hydroxylation site.

amino acids 253-265, 294-306, 335-347, 376-388, 417-423, 458-464,
540-546, 581-587

FIGURE 16

GGAGCCGCCCTGGGTGTCAGCGGCTGGCTCCGCGCACGCTCCGGCGTCGCGCAGCCTCG
GCACCTGCAGGTCGCGTCCCGGGCTGGCCCCCTGACTCCGTCGGGTTTTGTTCTGGGCTGA
CATGATTTCCCTCCGGGCCCCCTGGTGACCAACTTGTGCGGGTTTTGTTCTGGGCTGA
GTGCCCTCGGCCCTCGGGGCCAGCTGCAACTGCACTTGCCGCAACCGGTTGCAG
GCGGTGGAGGGAGGGGAAGTGGTGCCTCCAGCGTGGTACACCTTGACGGGAGGTGCTTC
ATCCCAGCCATGGGAGGTGCCCTTGTGATGTTCTCAAACAGAAAAGAAAAGGAGGATC
AGGTGTTGTCCTACATCAATGGGTACAACAAGCAAACCTGGAGTATCCTGGTCTACTCC
ATGCCCTCCGGAACCTGTCCCTCGGGCTGGAGGTCTCAGGAGAAAGACTCTGGCCCTA
CAGCTGCTCGTAATGTGCAAGACAAACAGGCAAATCTAGGGCACAGCATAAAACCT
TAGAACTCAATGTA(TGGTCTCCAGCTCCCATCCTGCCGTCTCCAGGGTGTGCCCCAT
GTGGGGCAAACGTGACCCCTGAGCTGCCACTCCAAGGAGTAAGCCCGTGTCCAATACCA
GTGGGATCGGCA(GCTTCCATCCTCCAGACTTCTTGACCCAGCATAGTGTATCCGTG
GGCTTTAACGCCACCAACCTTCTGCTTCCATGGCTGGAGTCTATGCTGCAAGGCCAC
AATGAGGTGGGACTGCCAATGTAATGTGACGCTGGAAGTGAGCACAGGGCTGGAGCTC
AGTGGTGTGGAGCTGGTGTGGTACCTGGCTGGACTGGGTTGCTGGCTGGCTGGTCC
TCTGTACCA(CCGCCGGGCAAGGCCCTGGAGGAGCAGCCAATGATATCAAGGAGGATGCC
ATTGCTCCCGAACCTGCCCTGGCCAAGAGCTCAGACACAATCTCCAAGAATGGGACCCCT
TCCCTGTCACTCCGACGACGCCCTCGGGCACCCATGGCCCTCCAGGCTGGTGCAT
TGACCCCCACGCCAGTCTCCAGCCAGGCCCTGCCCTACCAAGACTGCCACGACAGAT
GGGGCCACCCCTCAACCAATATCCCCCATCCCTGGTGGGTTCTCCCTGGCTTGAGCCG
CATGGGTGCTGCTGTGATGGTGCCTGCCAGAGTCAAGCTGGCTCTGGTATGATGAC
CCCACCACTCATTGGCTAAAGGATTGGGCTCTCCTCTATAAGGGTACCTCTAGCAC
AGAGGCCCTGAGTCATGGGAAAGAGTCACACTCTGACCCCTAGTACTCTGCCACCTCTC
TTTACTGTGGGAAACCATCTCAGTAAGACCTAAGTGTCCAGGAGACAGAAGGAGAAGAGGA
AGTGGATCTGGAAATTGGGAGGAGCCTCACCCACCCCTGACTCTCTTATGAAGGCCAGCTG
CTGAAATTAGCTACTCACCAAGAGTGAGGGCAGAGACTTCCAGTCAGTGAGTCTCCAGGC
CCCCCTGATCTGACCCACCCCTATCTAACACCACCCCTGGCTCCACTCCAGCTCCCTGT
ATTGATATAACCTGTCAAGCTGGCTGGTTAGGTTACTGGGGCAGAGGATAGGGAATCTC
TTATTAACATGAAATATGTGTTGTTTCAATTGCAAAATTAAAGATAACATAA
TGTTGTATGAAAAA

FIGURE 17

MISLPGPLVTNLLRFLFLGLSALAPPSRAQLQLHLPANRLQAVEGGEVVILPAWYTLHGEVSS
SQPWEVPFVMWFQKQKEKEDQVLSYINGVTTSKPGVSLVYSMPSRNLSLRLEGLQEKDSGPY
SCSVNVQDKQGKSRGHSIKTLELNVLVPPAPPSCRLQGVPHVGANVTLSCQSPRSKPAVQYQ
WDRQLPSFQTFFAPALD VIRGSLSLTNLSSMAGVYVCKAHNEVGTAQCNVTLLEVSTGPGAA
VVAGAVVGTTLVGLGLLAGLVLLYHRRGKALEEPANDI KEDAIAPRTLWPWKSSDTISKNGTL
SSVT SARALRPPHGP PRPGAL TPTPSLSSQALPS PRLPTTDGAHPQPISPIPGGVSSSGLSR
MGAVPV MVPAQS QAGSLV

Signal peptide:

amino acids 1-29

Transmembrane domain:

amino acids 245-267

N-glycosylation site.

amino acids 108-112, 169-173, 213-217, 236-240, 307-311

N-myristoylation site.

amino acids 90-96, 167-173, 220-226, 231-237, 252-258, 256-262,
262-268, 308-314, 363-369, 364-370

Prokaryotic membrane lipoprotein lipid attachment site.

amino acids 164-175

FIGURE 18

FIGURE 19

MKRPLLLVVFSTLLNCSYTQNCTKTPCLPNAKCEIRNGIEACYCNMGFSGNGVTI CEDDNEC
GNLTQSGENANCNT EGSYCCMVCVPGF RSSNQDRFTIN DGTVCI ENVNANCHLDNV CIAA
NINKLTKIRSIKEPVALLQEVYRNSVTDLSP TDITIYIEILAESSSL LGYKNNTISAKDTL
SNTLTFEVKTVNNFVQRDTFVWDKLSVN HRRTHLTKLMHTVEQATLRIQS FQKTTEFDT
NSTDIALKVFFFDSYNMKHIIHNMMDGYINIFPKRKAAYDSNGNVAVAFLYYKSIGPLLS
SSDNFLLKPKQNYD NSEEEERVISSVI SVSMSNPP TLYELEKITPTLSRKVTDYRSICAF
WNYS PDMNGWSSEGCELTYSNETHTSCR CNHLTHFAILMSSGPGSIGIKDYNILTRITQLG
IIISLICLAICIFTWFFFSEIQSTRTTIHKNLCCSLFLAEI VFLVGINTNTNKLFC SIIAGL
LHYFFLAFAFWMCIEGIHLLIVVGVIYNKGFLHKNFYIIFGYLSPAVVVGFSAALGYRYYGT
TKVCWLSTENNFIWSFIGPACLI IILVNLLA FGVI IYKVFRHTAGLKPEVSCFENIRSCARGA
LALLFLLGTTWIFGVLVHVVHASVVTAYLFTVSNAFQGMFIFLFLCVLSRKI QEEYYRLFKNV
PCCFGCLR

Signal peptide:

amino acids 1-19

Transmembrane domain:

amino acids 430-450, 465-486, 499-513, 535-549, 573-593, 619-636,
648-664

N-glycosylation site.

amino acids 15-19, 21-25, 64-68, 74-78, 127-131, 177-181,
188-192, 249-253, 381-385, 395-399

Glycosaminoglycan attachment site.

amino acids 49-53

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 360-364

Casein kinase II phosphorylation site.

amino acids 54-58, 68-72, 76-80, 94-98, 135-139, 150-154,
155-159, 161-165, 181-185, 190-194, 244-248, 310-314, 325-329,
346-350, 608-612

Tyrosine kinase phosphorylation site.

amino acids 36-44, 669-677, 670-678

N-myristoylation site.

amino acids 38-44, 50-56, 52-58, 80-86, 382-388, 388-394,
434-440, 480-486, 521-527

Aspartic acid and asparagine hydroxylation site.

amino acids 75-87

FIGURE 20

TGGAAACATATCCTCCCTCATATGAATATGGATGGAGACTACATAAATATTTCAAAGNG
AAAAGCCGGCATATGGATTCAATGGCAATGTTGCAGTTGCATTTTATATTATAAGAGTAT
TGGTCCCTTGCTTCATCATCGACAACCTCTTATTGAAACCTCAAATTATGATAATTCT
GAAGAGGGAGGAAAGAGTCATATCTCAGTAATTCAGTCTCAATGAGCTCAAACCCACCC
ATTATATGAACTTGAAAAATAAACATTACATTAAGTCATCGAAAGGTACAGATAGGTATA
GGAGGTCTATGTGGCATTTGGAATACTCACCTGATACCATGAATGGCAGCTGGTCTTCAGAG
GGCTGTGAGCTGACACTCAAATGAGACCCACACCTCATGCCGCTGTAATCACCTGACACA
TTTGCAATTGATGTCCTCTGGCCTTCCATTGGTATTAAAGATTATAATATTCTTACAA
GGATCACTCAACTAGGAATAATTATTCAGTGAATTGCTTGCCATATGCATTTCACCTTC
TGGTTCTTCAGTGAAATTCAAAGCACCAGGA

FIGURE 21

FIGURE 22

MRSGCVVVHVWILAGLWLAVAGRPLAFSDAGPHVHYGWGDPIRLRHLYTSGPHGLSSCFLRI
RADGVVDCARGQSAHSLLIEIKAVALRTVAIKGVHSVRYLCMGADGKMQQLQYSEEDCAFEET
EIRPDGYNVYRSEKHRLPVSLSAKQRQLYKNRGFLFLSHFLPMLPMVPEEPEDLRGHLESD
MFSSPLETD SMDPFGLVTGLEAVRSPSFEK

Signal peptide:

amino acids 1-22

Casein kinase II phosphorylation site.

amino acids 78-82, 116-120, 190-194, 204-208

N-myristoylation site.

amino acids 15-21, 54-60, 66-72, 201-207

Prokaryotic membrane lipoprotein lipid attachment site.

amino acids 48-59

FIGURE 23

CCCAGAAGTTCAAGGGCCCCGGCTCCTGCGCTCCTGCCGCCGGACCCCTGACCTCCTCA
GAGCAGCCGGCTGCCGCCGGAAAGATGGCGAGGAGGAGCCACC GCCCTCCCTGCTG
CTGCTGCCTACCTGGTGGTCGCCCTGGGCTATCATAGGCCCTATGGGTTTCTGCCAAAA
AGACCAACAAGTAGTCACAGCAGTAGAGTACCAAGAGGCTATTTAGCCTGCAAACCCAA
AGAAGACTGTTCCCTCCAGATTAGTGGAAGAAAACTGGGCTGGAGTGCTCCTTGCTAC
TATCAACAGACTCTTCAAGGTGATTTAAAAATCGAGCTGAGATGATAGATTCAATATCCG
GATCAAAATGTGACAAGAAGTGATGCCGGAAATATCGTTGTGAAGTTAGTGCCTCATCTG
AGCAAGGCCAAAACCTGGAAGAGGATAACAGTCAGTCTGGAAAGTATTAGTGGCTCCAGCAGT
CCATCATGTGAAGTACCCCTCTGCTCTGAGTGGAACGTGGTAGAGCTACGATGTCAAGA
CAAAGAAGGAATCCAGCTCTGAATACACATGGTTAACAGGATGGCATCCGTTGCTAGAAA
ATCCCAGACTTGGCTCCAAACCCACAGCTCATACACAATGAATAACAAAATGGAACT
CTGCAATTAAATCTGTTCCAAACTGGACACTGGAGAATATTCTGTGAAGGCCGCAATT
TGTGGATATCGCAGGTGCTGGAAACGAATGCAAGTAGATGATCTAACATAAGTGGCA
TCATAGCAGCCGTAGTAGTTGGCCTTAGTGATTTCCGTTGTGGCCTTGGTGATGCTAT
GCTCAGAGGAAAGGCTACTTTCAAAAGAAACCTCCCTCCAGAACAGTAATTCTCATCTAA
AGCCACGACAATGAGTGAAAATGTGCACTGGCTCACGCCCTGTAATCCACACTTGGAGG
CCGGCGGGCGGATCACGAGGTCAAGGAGTTCTAGACCAGTCTGGCAATATGGGAAACCC
CATCTCTACTAAAATACAAAATTAGCTGGCATGGTGGCATGTGCCTGCAGTTCCAGCTGC
TTGGGAGACAGGAGAATCACTTGAACCCGGGAGGCGGAGGTTGCACTGAGCTGAGATCACGC
CACTGCAGTCCAGCCTGGTAAACAGAGCAAGATTCCATCTAAAAAATAAAATAAATA
AATAAAATCTGGTTTACCTGTAGAATTCTTACAATAATAGCTTGATATTG

FIGURE 24

MARRSRHRLLLLLLRYLVVALGYHKAYGFSAPKDQQVVTAVEYQEAILACKTPKKT
WKKLGRSVSFVYYQQTTLQGDFKNRAEMIDFNIRIKNVTRSDAGKYRCEVSAPSEQGQNLEED
TVTLEVVLVAPAVPSCEVPSSALSGTVVELRCQDKEGNPAPEYTWFKD
GIRLLENPRLGSQST
NSSYTMNTKTGTLQFNTVSKLDTGEYSCEARNSVGYRRCPGKRMQVDDLNI
SISGIIAAVVVA
LVISVCGLGVCYAQRKGYSKETSFQKSNSSSKATTMSENVWLTPVIPALW
KAAAGGSRGQEF

Signal peptide:

amino acids 1-20

Transmembrane domain:

amino acids 130-144, 238-258

N-glycosylation site.

amino acids 98-102, 187-191, 236-240, 277-281

Casein kinase II phosphorylation site.

amino acids 39-43, 59-63, 100-104, 149-153, 205-209, 284-288

N-myristoylation site.

amino acids 182-188, 239-245, 255-261, 257-263, 305-311

Amidation site.

amino acids 226-230

FIGURE 25

GACATGGAGGGGGCTAGCACTGAAACTGCTTTCAAGACGAGGAAGAGGAGGAGAAGAG
AAAAGAGGAAGATGTGGGCAACATTATTTAACATGCTCCACAGCCGGACCTGGCAT
CATGCTGCTATTCCTGCAAATACTGAAGAACATGGGATTAAATATTACTCTAAATAA
ATGAATTACTCAATCCTCATGACCATCTATACATCTCCACCTTCAAAAGTACATCAATA
TTATATCATTAAGGAAATAGTAACCTTCTCTTCAATCGCTCCTATTGAACCTACTGCACTG
CAATTGTGCACTGGCACTTATTCTCATGTAAGAAAAAACTTTGTGGTTCTATGGCAATTCA
TTTGACAAATGCAGCATCTCCCTATCAATCGCTCCTATTGAACCTACTGCACTG
TGGAAATCCTTAAGGGGCCATTACATTCTGAAGAAGAACGTAAGATGAAGGACATGCCACT
CCGAATTCTATGTGCTACTTGGCCTAGTCACTACACTACAGTACAAGCTGTAGATAAAAAG
TGGATTGTCCACGGTTATGTACGTGAAATCAGGCCCTGGTTACACCCAGATCATTAT
ATGGAAGCATCTACAGTGGATTGTAACTGATTAGGTCTTTAACTTCCAGGCCAGATTGCC
AGCTAACACACAGATTCTCTCTCACAGATAACAAATATTGCAAAATTGAAATCTCCACAG
ACTTCCAGTAACACTTACTGCGCTGGATTATCTCAAAACAAATTATCTTCACTGCA
ATTAATGTAAAAAAAGATGCCCTAGCTCCTTCTGTGACTTAGAGGAAACAAACTTACTGA
ACTGCCCTGAAAATGCTGTGCGAACACTGAACTTACAAGAACACTCTATTAATCACA
TCTCTTCTCAATTCTACCTGGAGCCTTATGGCCTACATAATCTTCTCGACTTCATCTC
AATTCAAATAGATTGCGAGATGATCACACAGTAAGTGGTTGATGCTCTCCAATCTAGAGAT
TCTGATGATTGGGAAAATCCAATTATCAGAACTCAAAGACATGAACCTTAAGCCTCTTATCA
ATCTCGCAGCTGGTTAGCTGGTATAACACTCAGACAGATAACGCCCTGGTT
GGACTGAAAACCTAGAAAGCATCTTCTTACGATAACAGGCTTATTAAAGTACCCCATGT
TGCCTCTCAAAAAGTGTAAATCTCAAATTGGATCTAAATAAAAATCTTATTAATAGAA
TACGAAGGGGTGATTAGCAATATGCTACACTTAAAGAGATTGGGATAAATAATGCGCT
GAGCTGATTTCATCGATAGTCTGCTGTGGATAACCTGCAGATTAAAGAAAATAGAAC
TACTAACACCTAGATTGTCTTACATTACCCCCATGCTTTTCACTGACTCCCCAAGCTGG
AATCACTCATGCTGAAACAGCAATGCTCTCACTGGCCCTGTACCATGGTACATTGAGTC
CCTAACCTCAAGGAAATCAGCATACACAGTAAACCCATCAGGTGTGACTGTGCTATCCGGT
GATGAACATGCAACAAAACCAACTTCGATTCTGGACAGGCGAGATTCACTGTTTGC
CACCTGAATTCCAAGGTCAAAGTGTGCGCAAGTGCAATTCCAGGGACATGATGGAAATTG
CTCCCTCTTATAGCTCTGAGGCTTCTCTTAATCTAAATGTAAGGGTCTGGAGCTATGT
TCTCTTCACTGTAGAGCTACTGCGAGAACACCAGCTGAAATCTACTGGATAACACCTTCTG
GTCAAAACTCTTGCCTAATACCCATGACAGACAAGTCTATGTCATTCTGAGGGAAACACTA
GATAAAATGGCTAACTCCCAAGAAGGGGTTTATATACTTGTATAGCAACTAACCTAGT
TGGCGCTGACTTGAAGTCTGTATGATAAAGTGGATGGATCTTTCCACAGATAACATG
GCTCTTGAATATAAAAAGAGATACTCAGGCCAATTCTGGGTTCTGGTCTGGAAAGCA
AGTTCTAAAATTCTCAAATCTAGTGTAAATGGACAGCCTTGTCAAGACTGAAAATTCTCA
TGCTCGCAGGAGTGTGCAATACCATCTGTGTCAGGTATATACTTACTCATCTGAATC
CATCACTGAGATAAAAATTGTATTGATATTCCACCATCTATCAGAAAACAGAAAAAAA
TGTGTAAATGTCAACCAAGGTTGCAACCTGATCAAAAGAGTATGAAAAGAATAATAC
CACACACTTATGGCCTGCTGGAGGCCCTGTGGGATTATTGGTGTGATATGTCCTTATCA
GCTGCCCTCTCCAGAAAATGAAGTGTGGACAGCAGTGTGAGGAATTACTTACAG
AAACCAACCTTGTGCTTACGAGCTTATCTCTCTGTATAATCTCTGGAAAGCAGAAA
AGAAAAAAAGTACATCACTGAAAGTAAAGCACTGTTAGGTTACACAAATAATGTCCT
AAAAACCAAGGAAACCTACTCCAAAATGAAC

FIGURE 26

MKDMPLRIHVLLGLAITTLVQAVDKKVDCPRLCTCEIRPWFTPRTSIYMEASTVDCNDLGLLT
FPARLPANTQILLQTNIAKIEYSTDFPVNLTGDLDSQNLLSSVTNINVKKMPQLLSVYLE
ENKLTTELPEKCLSESLNLQELYINHNLLSTISPGAFIGLHNLRLHLNSRQLQMINSKWFDA
LENLEILMIGENPIIIRIKDMNFKEPLINLRSVIAGINLTEIPDNAVGLENLESISFYDNRL
IKVPHVALQKVVNVLKFLLDNKPNPINRIRRGDFSNMLHILKELGINNNPELISIDS LAVDNLPD
LRKIEATNNPRLSYIHPNAFFRLPKLESLMNNSNALSA LYHTIESLPNLKEISIHSNPIRC
DCVIRWMNNMKTNIRFMEPDSLFCVDPPEFQGQNVRQVHFDRDMMEICLPLIAPESFPSNLNV
EAGSYVSFHCRAATEPOPEIYWITPSGQKLLPNTLTDKFYVHSEGTLIDINGVTFKEGGLYTC
IATNLVGADLKSVMIKVDGSFPQDNNGSLNIKIRDIQANSVLVSKASSKILKSSVKWTAFV
KTENSHAAQSARIPSDVVKVYNLTHLNPSTEYKICIDIPTIYQKNRKCVNVTTKGLHPDQKE
YEKNNNTTLMACLGGLLGIIGVICLISCLSPEMNCDDGHHSYVRNYLQKPTFALGELYPPLIN
LWEAGKEKSTS LKV KATVIGLPTNMS

Signal sequence:

amino acids 1-22

Transmembrane domain:

amino acids 633-650

N-glycosylation site.

amino acids 93-97, 103-107, 223-227, 382-386, 522-526, 579-583,
608-612, 624-628, 625-629

Casein kinase II phosphorylation site.

amino acids 51-55, 95-99, 242-246, 468-472, 487-491

Tyrosine kinase phosphorylation site.

amino acids 570-579

N-myristoylation site.

amino acids 13-19, 96-102, 158-164, 221-227, 352-358, 437-443,
491-497, 492-498, 634-640, 702-708

Cell attachment sequence.

amino acids 277-280

FIGURE 27

GGCCGGGACTGGCGAAGGTGCCAAGCAAGGAAAGAAATAATGAAGAGACACATGTGTTAG
CTGCAGCCTTTGAAACACGCAAGAAGGAATCAATAGTGTGGACAGGGCTGGAACCTTAC
CACGCTTGGAGTAGATGAGGAATGGGCTGTTGATTATGCTGACATTCCAGCATGAATCT
GGTAGACCTGTGGTTAACCGTCCCCTCTCCATGTGTCCTCCTACAAGGTTGTTCTTA
TGATACTGTGCTTCATTCTGCCAGTATGTGCCCCAAGGGCTGCTTGTCTTCCCTCTGGG
GGTTAAATGTCACTGTAGCAATGCAAATCTCAAGGAAATACCTAGAGATCTTCCCTCTGA
AACAGTCTTACTGTATCTGGACTCCAATCAGATCACATCTATTCCAATGAAATTTTAAGG
ACCTCCATCAACTGAGAGTTCTCAACCTGTCCAAAATGGCATTGAGTTATCGATGAGCAT
GCCTTCAAAGGAGTAGCTGAAACCTTGAGACTCTGGACTTGTCCGACAATGGATTCAAAG
TGTGCAACAAATGCCCTCAATAACCTGAAGGCCAGGGCCAGAATTGCCAACACCCCTGGC
ACTGCGACTGTACTCTACAGCAAGTTCTGAGGAGCATGGCGTCCAAATCATGAGACAGCCAC
AACGTGATCTGAAAACGTCGTTGGATGAAACATGCTGGCAGACCATTCTCAATGTC
CAACGACGCTGACCTTGTAAACCTCCCTAAAAAAACTACCGATTATGCCATGCTGGTCACCA
TGTTGGCTGGTTCACTATGGTATCTCATATGTGGTATATTATGTGAGGCAAATCAGGAG
GATGCCCGGAGACACCTCGAATACTTGAATCCCTGCCAACGAGGAGAAAGCAGATGA
ACCTGATGATATTAGCACTGTGGTATAGTGTCCAAACTGACTGTCAATTGAGAAAGAAAGAAA
GTAGTTGCGATTGCAAGTAGAAATAAGTGGTTACTCTCCCATCCATTGTAACACATTGAA
ACTTGTATTTCACTTTTTTGAAATTATGCCACTGCTGAACCTTAACAAACACTACAACA
TAAATAATTGAGTTAGGTGATCCACCCCTTAATTGTACCCCCGATGGTATATTCTGAGT
AAGCTACTATCTGAACATTAGTTAGATCCATCTCACTATTAAATAATGAAATTATTTTT
AATTAAAGCAAAAGCTTAACCTTGAAACCATGGAAAAAAAAAAAAAAACA

FIGURE 28

MNLVDLWLTRSLSMCLLQSFLMILCFHSASMC PKGCLCSSGGLNVTCSNANLKEIPRDL
PPETVLLYLDNSQITSIPNEIFKDLHQLRVLNLSKNGIEFIDEHAFKGVAETLQTLDSLSDNR
IQSVHKNAFNNLKARARIANPWHCDCTLQQVLRSMASNHETAHNVIKTSVLDEHAGRPF
NAANDADLCNLPKKTTDYAMLVTMFGWFTMVISYVVYYVRQNQEDARRHLEYLKS LPSRQKK
ADEPDDDISTVV

Signal sequence:

amino acids 1-33

Transmembrane domain:

amino acids 205-220

N-glycosylation site.

amino acids 47-51, 94-98

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 199-203

Casein kinase II phosphorylation site.

amino acids 162-166, 175-179

N-myristoylation site.

amino acids 37-43, 45-51, 110-116

FIGURE 29

ACCGAGCCGAGCGGACCGAAGGCAGCAGGAGATGCAAGGTGAGCAAGGGATGCTGGGGGG
GGCCTGAGGAGCATGCCAGCCCCCTCTTGGCCCTCTGAGGAGCATCTCCCTCTGCTGCT
GGCTCTAGTCCTGCTCAGGCTCGGCCAACGGGCTGGCCCGGCGCTGCGATGCTCCGCCAG
ACCGCGCTGTGCTGTGCCACCGCAAGTGCTTGTGGCAGTCCCCGAGGGCATCCCCACCGAG
ACCGCCTGCTGACCTAGGCAAAACGGCATCAAACGCTCAACCAGGACGAGATTGCCAG
CTTCCCAGGACTGGAGGAGCTGGAGCTCAACGAGAACATCTGAGCCCGTGGAGCCCGG
CTTCAACAAACCTCTTCAACCTCCGGACGCTGGGCTCTCCGGCACAAACCCCTGAAGCTCATC
CGGCTAGGCGTCTCACTGGCCTCAGCAACCTGACCAAGCAGGACATCAGCGAAACAGAT
CGTTATCTCTGACTGACTCATGTTCTGGACCTGTACAACCTCTGAAGTACTGGAGGTG
AAATGACCTCTGCTCATCTCTACCGGCCCTCATGGGCTCTAACAGCTGGAGCAGCTG
ACGCTGGAGAAATGCAACCTGACCTCCATCCCCACCGAGGGCTGTCCACCTGACGGCCT
CATCGCTCTGAGGCTCCGGCACCTCAACATCAATGCCATCCGGACTACTCTCTCAAGAGGC
TGTAACGACTCAAGGCTTGGAGATCTCCACTGGCCCTACTTGGACCATGACACCCAA
TGCTCTACGGCCTCAACTCTGACGCTCTCTGTCATCACACACTGCAATCTGACCGCTGTG
CTACCTGGCCGTCGCCACCTAGTCTATCTCGCTTCTCAACCTCTCTACAAACCCATCA
GCACCATCTGGGGCTCATGGTGTGACGCTGTCGGCTGCGAGGATTCAGCTGGTGGG
GGGCAGCTGGCGTGGTGGAGGCCATTGCTTCCGGCCCTCAACTCTGGCGCTGCTCAA
TGTCTCTGGCAACCAGCTGACCAACTGGAGGAATCAGTCTTCACTCGGTGGCAACCTGG
AGACACACTCTGGGACTCTAACCGGCTGGCCCTGCGACTGCGCTCTGGTGTGGTCT
CCGGCGTGGCGCTCAACTCTAACCGGCAAGCAGGCCACCTGGCCGACCGGGAGTTGGTCT
GGGAAGGGAGTTCAAGGACTTCCCTGATGTGCTACTGCCAACTACTCTACCTGGCCGGCG
CCGGCAGTCTGGGAGCCGAAAGGGCACGAGTGGTGTGGACGAGGGCACACGGTGTGG
GTGTGGGGGGCATGGCCAGGGCCAGGGCCGGCCCTACCTCTGGCTCTACCCGGAAAGACCT
GGTCTCAGGCAAGAGCAATGGGCGCTCACAGTCTTCCCTGATGGCACGCTGGAGGTG
ACGCCAGGTACAGGAAACGGCAGTACCTGTGATCGCGCCAAACGGGGCGGCAACGAC
TCCATGGCCGCCACCTGATGTGCGACTGCGCTACTGCCAGCTGGCCCATGCCCACAA
GACCTCTGGCTTCTCATCTAACACAGGCCGGGGAGGGAGGCCAACAGCACCCTGGCCT
TGCCCTTCCCCCTGACATCAAGACCTCATCATGCCACCCATGGGCTTATCTCTTCT
CTGGGGCTGCTCTCTCTGGCTGTGCTGTTCTCTGGAGGCCGGCAAGGGCACAC
AAAGCACACATCGAGATCTGGAGTATGCTGGCCCGAAGTCGAGCAGGGCATCAGCTCCGG
ACGGCCGGCGCAAGTTCAACATGAAGATGATAATGAGGGCGGGGGGGAGGGACCCCCG
ACGGCCGGGGAGGGGAAGGGGCTGGTCCGACCTCTGCTACTCTCCAGTCTTCCACCTC
CTCCCTACCTCTACACAGTCTCTTCTCTCCGGCTCTGGCTGCTGTCCTGGCC
CCAGGCTTACACCTGGCTCTTCTACAGGACCTGAGGACAGGACAGGGCACCTGGG
CCTACACAGGGGATTGACAGACTGGAGTTGAAAGCCAGAACGCCAACCGCAGAGTCA
TAATTCAATAAAAAGTACAGAATTCTCTGTAACATTGGGTTCAATAATTGATGATT
TATGAAAATTCTGAAAATAAAAAGGAAAAAACTAAAACAAAAAAACAAAAAA

FIGURE 30

MQVSKRMLLAGGVRSRMPSPLLACWQPILLLVLSGSATGCPRCECSAQDRAVLCHRKCF
VAVPEGIPTETRLLDLGKNRIKTLNQDEFASFPHLEELELENENIVSAVEPGAFNNLFNLRTL
GLRSNRKLIPLGVFTGLSNLTKQDISENKIVILLDYMFQDLYNLKSLEVGDNDLVYISHRA
FSGLNSLEQLTLEKCNLTSIPTEALSHLHGLIVLRLRHNLNAIRDYSFKRLYRLKVLEISH
WPYLDTMTPNCLYGLNLTSLSITHCNLTAVELYLAVRHLVYLRFLNLSYNPISTIEGSMIHEL
LRLQEIQLVGGQLAVVEPYAFRGLNYLRVLNVSGNQLTTLLEESVFHSVGNLETLLIDSNPLA
CDCRLLWVFRRWRLLNFNRQOPTCATPEFVQGKEFKDFPDVLLPNYFTCRRARIRDRKAQOV
FVDEGHTVQFVCRADGDPPAIIWLSPRKHIVSAKSNGRLTVFPDGTLLEVRYAQVQDNQTYL
CIAANAGGNDSMPAHLHVRSYSPDWPHQPNKTFAFISNQPGEGEANSTRATVPFPFDIKTLI
IATTMGFISFLGVVLFCLVLLFLWSRGKGNTKHNIELEYVPRKSDAGISSADAPRKFNMKMI

Signal sequence:

amino acids 1-41

(P) Proline (A) Alanine (G) Glycine (S) Serine (T) Threonine
(C) Cysteine (Y) Tyrosine (W) Tryptophan (F) Phenylalanine
(I) Isoleucine (V) Valine (D) Aspartic acid (E) Glutamic acid
(N) Asparagine (Q) Glutamine (H) Histidine (K) Lysine (R) Arginine
(L) Leucine (M) Methionine (U) Cysteine (B) Alanine or Threonine
(X) Unknown (O) Oxidized cysteine (D*) Aspartic acid (E*) Glutamic acid

Transmembrane domain:

amino acids 556-578

N-glycosylation site.

amino acids 144-148, 202-206, 264-268, 274-278, 293-297, 341-345,
492-496, 505-509, 526-530, 542-546

Casein kinase II phosphorylation site.

amino acids 49-53, 108-112, 146-150, 300-304, 348-352, 349-353,
607-611

Tyrosine kinase phosphorylation site.

amino acids 590-598

N-myristoylation site.

amino acids 10-16, 32-38, 37-43, 113-119, 125-131, 137-143,
262-268, 320-326, 344-350, 359-365, 493-499, 503-509, 605-611

Prokaryotic membrane lipoprotein lipid attachment site.

amino acids 32-43

FIGURE 31

CCCCACGCGTCCGCACCTCGGCCCCGGGCTCGAAGCGGTCTGGGGGCCCTTCGGTCAAC
ATCGTAGTCACCCCCCTCCCCATCCCCAGCCCCCGGGGATTCAAGGCTCGCCAGGCCAGCC
AGGGAGGCCGGGAAGCGCGATGGGGGCCAGCGCCTCGCTCTGCTCTGCTCTG
TGTTCGCTGCTGCTGGGCCCGCGGGCCAACCTCTCCAGGACGACAGCCAGCCCTGG
ACATCTGATGAAACAGTGGTGCTGGTGACCGTGGTCAAGTCCAAGTGAAAGATCA
CGAGGACTCATCCCTGCAATGGCTAACCTGCTCAGCAGACTCTACTTTGGGGAGAAGA
GAGCCCTTGAGATAATCGAATTAGCTGGTACCTCTACGCCACAGAGCTCAGCATCAGC
ATCAGCAATGTGCCCTGCCAGACAGGGCAGTACACTGCTCAATCTTCACTATGCCGT
GCCAAGTCCAAGTCCCCTGCACTGTGCTAGGAATTCCACAGAACGCCATCATCACTGGT
ATAAAATCTCATTACGGGAAAAGACACAGGCCCTAAACTGTCACTTCTGGGAGCAAG
CCTGCAGCCCGGCTCACCTGGAGAAAGGGTGACCAAGAACCTCACGGAGAACCAACCGC
ACAGGAAGATCCAATGGTAAACCTCACTGTCAGCAGCTGGTACATTCCAGGTTACCC
GGGAGGATGATGGGCGAGCATCGTGTGCTGTGAACCATGAATCTAAAGGGAGCTGAC
AGATCCACCTCTCAACGATTAGAGTTTATACACACCAACTGCGATGATTAGGCCAGACCC
TCCCCATCTCGTGGGGCAGAAGCTGTTCTACACTGTGAGGGTGCAGGCAATCCAGTCC
CCCAGCAGTACCTATGGGAGAAGGGCAGTGTGCCACCCCTGAAGATGACCCAGGAGAGT
GCCCTGTACTTCCCTTCTCAACAGAGTGAAGTGGCACCTACGGTGCACAGCCACAG
CAACATGGCAGCTACAAGGCTACTACACCTCAATGTTAATGACCCAGTCGGTGCCT
CCTCCTCCAGCACCTACCAAGGCCATCATGGGGATCGGGCTTCATTGCTTCCGTG
CTCATCATGCTCATCTCCGGCCACTACTGTGATCCGGCACAAAGGAACCTACCTGACACA
TGAGGCAAAGGCTCCGACGATGCTCCAGCAGCGGACACGGCCATCATCAATGCGAGAAGGCG
GGCAGTCAGGGAGGAGCACAAGAAGGAATTTCATCTAGAGGCGCTGCCACTTCTGC
GCCCCCAAGGGCCCTGGGGACTGCTGGGCCGTACCAACCCGACTTGTACAGACAA
CCGCAGGCCGCCCTCCGCTGCTCCCCAGGCCACCCACCCCTGTACAGAAATGCTGC
TTGGGTGGGTTTGTACTCGGTTGGAAATGGGGAGGGAGGAGGGGGGGGGGGAGGG
TTGCCCTCAGCCCTTCCGTCGGCTCTGCAATTGGGTTATTATTATTTGTAACATCC
CAAATCAAATCTGTCCTCAGGCTGGAGAGGCAGGAGCCCTGGGTGAGAAAAGCAAAAACAA
ACAAAAAAACA

FIGURE 32

MGAPAASLLLLLFFACCWAPGGANLSQDDSQWPWSDETVVAGGTVLKCQVKDHEDSSLQW
SNPAQQTLYFGEKRALRDNRIQLVTSTPHELSIISNVALADEGEYTCISIFTMPVRTAKSLV
TVLGIPQKPIITGYKSSLREKDATTLCQSSGSKPAARLTWRKGQELHGEPTRIQEDPNGK
TFTVSSSVTQVTREDDGASIVCSVNHESLKGADRSTSQRIEVLYTPTAMIRPDPPHPREGQ
KLLLHCEGRGNPVPQQYLWEKEGSVPPLKMTQESALIFPFLNKSDSGTYGCTATSNMGSYKA
YYTLNVNDPSPVPSSSSTYHAIIGGIVAFIVFLLLIMLIFLGHYLIIRHKGTYLTHEAKGSDD
APADATIAAEGGQSGGDDKKEYFI

Signal sequence:

amino acids 1-20

Transmembrane domain:

amino acids 331-352

N-glycosylation site.

amino acids 25-29, 290-294

Casein kinase II phosphorylation site.

amino acids 27-31, 35-39, 89-93, 141-145, 199-203, 388-392

N-myristoylation site.

amino acids 2-8, 23-29, 156-162, 218-224, 295-301, 298-304,
306-310, 334-340, 360-364, 385-389, 386-390

Prokaryotic membrane lipoprotein lipid attachment site.

amino acids 7-18

FIGURE 33

GGGGGTTAGGGAGGAAGGAATCCACCCCCACCCCCCCTTAAACCCCTTTCTTCCTTCTGG
CTCGGACATTGGAGCACTAAATGAATTGTGTCGTGCGAGCAGGATGGTCGCTG
TTACTTTGTGATGAGATCGGGGATGAATTGCTGCCTTAAAAATGCTGCTTGGATTCTGTT
GCTGGAGACCTCTCTTTGCGCTGGAAAAGCTTACAGGGGACGTTGCAAAGAGAAGA
TCTGTTCCCTGCAATGAGATAGAAGGGGACCTAACAGTAGACTGTGAAAAAAAGGGCTTCACA
AGCTGCAAGCGTTTCACTGCCCGACTTCCAGTTTACCATTTATTCTGCATGCCAATT
CCTCACTCGAATTTCCTTAATGAGTTGCTAACCTTTATAATGCGGTTAGTTGCACATGG
AAAACAATGGCTTGCAATGCTTCCGGGGCTTTCTGGGGCTGCAGCTGGTGAAAAGG
CTGCACATCAACAAACAAGATCAAGTCTTCTGAAAGCAGACTTTCTGGGGCTGGACGA
TCTGGAATATCTCAGGCTGATTTTAATTATAGCAGATATAGACCCGGGGCTTCAGG
ACTGAAACAGCTGGAGGTGCTCAATTAAATGACAATCTCATCAGCACCTACCTGCAAC
GTGTTCCAGTGTGCCCCATCACCCACCTGCCACCTCGGGAACAGGGCTGAAAACAGCTG
CTATGAGGAGCTTGGAGCAATTCCCTGGTATTGCGGAGATCTCTGCTAGAGGATAACCTT
GGGACTGCACTGTGATCTGCTCCCTGAAAGAATGGCTGAAAACATTCCAAGAATGCC
CTGATCGGCCAGGGCTCTGCAAGGCCCCACAGACTGCAGGGTAAAGACCTCAATGAAAC
CACCGAACAGGACTTGTGCTTGTGAAAGAACCTTGGACTGGAATTCTGCTCCGGGCCCTG
CCCAAGAAGAGACCTTGTCTGGACCCCTGCCAACACTCTTCAAGACAAATGGGCAAGAG
GATCATGCCACACCAGGGTCTGCTCCAAACGGAGGTACAAAGATCCAGGCAACTGGCAGAT
CAAATCAGACCACAGCAGCAGTGCAGGGTAGCTCAGGGAACAAACCCCTTAGCTGA
GTTACCTGCCCCGGGGCTGAGCTGCCACATCCCAGGGTGGGTTAAAGATGAAAC
TGCACAAACAGGAACGTGAGCAGCTGGCTGATTGAGGCCAAGCTCTAACGTGCAAGGA
GCTTTCTACAGAATAACAAGATCCACAGCATCGCAATTGCACTTTGTGATTACAAGA
ACCTCATTCTGTGATCTGGGACAACAATACATCGCTACTGCTAGAGAACACACTTCAAG
AACCTTTGGACTCAGGGCTATACATGGATAGCAATTACCTGGACACCGCTGCTCCGGGA
GAAATTCCGGGGCTGCAAAACCTAGAGTACCTGCAAGCTGGAGTACAACGCTATCCAGCTA
TCCCTCCGGGCACTTCAATGCCATGCCAAACTGGAGGACTCTCATTCTCAACAAACACTG
CTGAGGTCTCTGCCATGGACTGTTGCTGCTGGGTTCTGCTCTAAACTCAGGCTGCAAA
CAATTACTCATGATCACCTCCGGTGGCAGGGGCTGGACAGGTTAACCTCATCATCAGA
TAGACCTCACGGAAACCCCTGGGGAGTGTCTCTGCAAAATGTGCTCTTCAAGCAGTGGGA
GAACGCTGGGTTCTCGGAAGCTGTGATGGCAGGACCTCAAGTGTGAGACGCCGGTGAACCTT
TAGAAAGGATTTCATGCTCTCTCAATGACAGAGATCTGCCCTCAGCTGTACGCTAGGATCT
CGCCCACTGTTACCTCCGGCACAAGTAAAACAGCACTGGGTTGGCGAGACCGGGACGCACTC
AACTCTACCTAGACACCAGCAGGGTCTCGGCTGTTGGTCTGGCTGCTGCTGTT
GTTGTACCTCCGCTTCACGGTGGCATGCTGTTTATCCTGAGGGAAACGAAAGC
GGTCCAAGAGACAGATGCCAACTCCTCCGGCTCGAGATAATTCCCTACAGACAGTCTG
GACTCTTCTACTGGCACAATGGGCCCTAACAGCAGATGGGGCCACAGAGTGTATGACTG
TGGCTCTCACTCGCTCTGCAACTAGACCCCAACCCCAATAGGGGAGGGCAGAGGGAAAGGCG
ATACATCCTCCCCACCGCAGGACCCGGGGCTGGAGGGGCTGTAACCAATCCCCCG
CCATCAGGCTGGATGGGATAAGTAGATAATAACTGTGAGCTGCCACAAACGAAAGGGCT
GACCCCTTACTTAGCTCCCTCCTGGAAACAAAGAGCAGACTGTGGAGAGCTGGGAGAGCGCA
GCCAGCTGCTCTTGCTGAGAGCCCCCTTGCAGAGAAAGGGCAGCACGACCCCTGCTGGAG
AACTGACAGTGCCTCGCCCTGGGGGGGGCTGTGGGGTGGATGCCGGTTCTATAC
ATATATACATATCCACATCTATAGAGAGATAGATATCTTCTGGATGGATTAG
CCCCGGTGTGGCTCCCTGGCTACGCAAGGGATGCCAGTGCACGAAGGCATGAATGTAT
TGTAATAAGTAACCTTGACTCTGAC

FIGURE 34

MLLWILLLETSLCFAAGNVGTGVCKEKICSCNEIEGDLHVDCCEKKGFTSLQRFTA
PLFHLGNSLTRLFPNEFANFYNAVSLHMENNGLHEIVPGAPFLQLVLKR
TFLGLDDLEYLQADFNLLRIDPGAFQDLNKLEVILNDNLISTLPANVFQYVP
NRLKTLPLYEEVLEQIPGIAEILLEDPWDCDLSLKEWLENIPKNALIGRV
GKDLNETTEQDLCPLKRNVRDVSSLPAPPQAQETTFAPGPLPTPFKT
NQEDHATPGSAPNGGTKIPGNWQIKIRPTAAIATGSSRNKP
LANSLPCPCGGCSDHIPSGLKMNCNNRNVSSLADLKPK
KLSNVQELFLRDNIKHSIRKSHFVDYKNLILLLDGNNNIAT
AVENNTFKNLLDLRWLYMDSNYLDLTSREKFAGLQNLEYLN
VEYNAIQLILPCTFNAMPKLRILILNNNNLLRS
LPVDFVAGVSLSKSLHNYYFMYLPVAGVLDQTSIIQIDLHG
NPWBCSCTIVPFKQWAERLGSEVLMSDLKC
ETPVNFFRKDFMLLSNDIECPQLYARISPTLTS
HKSKNSTGAETGTHSN
SYLDTSRVSI
VPGLLL
V
F
V
T
S
A
F
T
V
V
G
M
L
V
F
I
L
R
N
R
K
R
S
K
R
R
D
A
N
S
A
S
E
I
N
S
L
Q
T
V
C
D
S
S
Y
W
H
N
G
P
Y
N
A
D
G
A
H
R
V
Y
D
C
G
S
H
S
L
S
D

Signal sequence:

amino acids 1-15

Transmembrane domain:

amino acids 618-638

N-glycosylation site.

amino acids 18-22, 253-257, 363-367, 416-420, 595-599, 655-659

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 122-126, 646-650

Casein kinase II phosphorylation site.

amino acids 30-34, 180-184, 222-226, 256-260, 366-370, 573-577,
608-612, 657-661, 666-670, 693-697

N-myristoylation site.

amino acids 17-23, 67-73, 100-106, 302-308, 328-334, 343-349,
354-360, 465-471, 493-499, 598-604, 603-609

Prokaryotic membrane lipoprotein lipid attachment site.

amino acids 337-348

FIGURE 35

AGTCGACTGCGTCCCTGTACCGGGCAGCTGTGTTCTGACCCCCAGAAATAACTCAGGGC
TGACCGGGCCTGGCACCGCTCCGACACATTCTGTGCGGGCTAAGGGAAACTGTGGC
CGCTGGGCCCGGGGGGATTCTTGGCAGTGGGGGTCGTCGGGAGGGCGAGGGGG
AAGGGGGGGGAAACGGGTTGGGAAGGCCAGCTGTAGAGGGCGGTGACCGCGCTCAGAACAC
AGCTCTGCGTCTCGAGCGGGACAGATCCAAGTGGCACTGGGAGAGCTCTGCGTGC
AGAATGAGCCGGCTTCGCCCCCTGTGCGCTCTGGCAGGGCTCTGGCCGGGGCTCAG
CGCGGAACACCCACTGCGCACCGTGCTGGCTCGCCCTCGGGGCTGCTACAGCTGC
ACCACGCTACCATGAACGGCAGGGCGCAGGGCCTGCATCCCTCGAGGTGGGGCGCTC
AGCACCGTGGTGCAGGGCGCCAGCTGCGCTGTGCTCGCCTCTGCGGGCAGGCCAGG
GCCCGGAGGGGGCTCAAAGACCTGCTGTTCTGGTGCACACTGGAGGGCAGGGTTCCACT
GCACCTGGAGAACAGGCTTTGCGGGGTTCTCTGTGGCTGTCTCCGACCCCGGGCTC
GAAAGCGCACCGCTGAGTGGTGGAGGAGCCCCAACCGCTCTGCAACCGCGGGAGATGCGC
GGTACTCAGGCCACCGGCTGGGGTGGGGCTGAGGCCAGGCTGAAGGAGATGCGATGCCACCTGC
GCCCAACGGTACCTGTGCAAGTACCAAGTTGAGGTCTTGTGCTCTGCGCGCCCCGGG
GCCGCTCTAACTTGAAGCTATCGCGCCCTTCAGCTGACAGGCCGCTCTGGACTTCAG
TCCACCTGGGACCGAGCTGAGTGGCTCTGCGGGGACAGCTCCGATCTCAGTTACTGCA
TCGCGGACAAATCGCGCTCGCTGGGACAACACTCTCGGGCGATGTGTTGTGCTCTGGCCC
GGGAGGTACCTCGTGGCAAATGCGCAGAGCTCCTAACTGCTAGACGACTTGGGAGG
CTTGCCTGGAATGTGTCAGGGCTTCGAGCTGGGAAGGAGACGGCCGCTTGTGACCA
GTGGGGAGGACAGCCGACCCCTGGGGGACGGGGTCCCACAGGCCGCCGGCCACT
GCAACCAAGCCCCGTGCCAGAGAACATGGCAATCAGGTGACGAGAACGCTGGGAGAGAC
ACCAACTTGTCCCTGAACAAGAACATTCTAGTAACATTCTGAGATTCTCGATGGGAT
CACAGAGCACGATGCTACCTCTCAAATGCTCTCAAGGCCAGTCAAAGGCCACTATCACC
CCATCAGGGAGCGTGAATTCCAAGTTAACTCTAGCTTCTCTGCCACTCTCAGGCTT
CGACTCTCTCTGCGCTGGTCTTCATATTGTGAGCACAGCAGTAGTAGTGTGTTGTGATCT
TGACCATGACAGTACTGGGGCTTGTCAAGCTGCTGCTTCAAGGAAAGCCCCCTTCCAGCCA
AGGAAGGAGTCTATGGGGCCGGGGCTGGAGAGTGTACTCTGAGCCCCTGCTTTGGGCTC
CAGTTGCAACATTGACAAAACATGGGGTGAAGTGGGACTGTAATCTGCGGGACAGAG
CAGAGGGTGCCTGCTGGGGAGTCCCCCTTGGCTTAGTGTGATGCA**TAGGAAACAGGGGA**
CATGGGCACTCTGTGAACAGTTTTCACTTTGATGAAACGGGGAAACCAAGAGGAACCTAC
TTGTGTAATGACAATTCTGAGAAATCCCCCTTCTCTAAATTCTTACTCCACTGAG
GAGCTAAATCAGAACCTGCACACTCCTCTGATGATGAGAGGAAGTGGAGTGCCTTCTAGGA
TGGTGTACTGGGGACGGGTAGTGGCTGGGGAGAGATTTCTTATGTTTATTCGGAGAA
TTGGAGAAGTGAATTGAACTTTCAAGACATTGGAAACAAATGAAACACAATATAATTACA
TTAAAAAAATAATTCTACCAAAATGGAAAGGAATGTTCTATGTTGTTCAAGCTAGGAGTAT
ATTGGTTCGAAATCCCAGGGAAAAAAATAAAAATTAAGGATTGTTGAT

FIGURE 36

MRPAPALCLLWQALWPAGPGGGEHPTADRAGCSASGACYSLHHATMKRQAAEAEACILRGGALS
TVRAGAELRAVLALLRAGPGPGGGSKDLLFWVALERRRSHTLENEPLRGFSWLSSDPGGLE
SDTLQWVEEPQRSCSTARRCAVLQATGGVEPAGWKEMRCHLRLANGYLCKYQFEVLCFAPRPGA
ASNLSYRAPFQLHSAALDFSPPGTEVSALCRGQLPISVTCIADEIGARWDKLSGDVLCPCPG
RYLRAGKCAELPNCLDDLGFACECATGFEIYGKDGRSCVTSGEGQPTLGGTGVPTRRPPATA
TSPVPQRTWPIRVDEKLGETPLVPEQDNSVTSIPEIPRWGSQSTMSTLQMSLQAESKATITP
SGVISISKFNNTSSATPQAFDSSSAVVIFVSTAVVVLVILMTVLGLVLCFHESPSSQPR
KESMGPPGLESDPEPAALGSSSAHCTNNGVKGDCDLRDRAEGALLAESPLGSSDA

Signal sequence:

amino acids 1-16

Transmembrane domain:

amino acids 399-418

N-glycosylation site.

amino acids 189-193, 381-385

Glycosaminoglycan attachment site.

amino acids 289-293

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 98-102, 434-438

Casein kinase II phosphorylation site.

amino acids 275-279, 288-292, 342-346, 445-449

N-myristoylation site.

amino acids 30-36, 35-41, 58-64, 59-65, 121-127, 151-157,
185-191, 209-215, 267-273, 350-356, 374-380, 453-459, 463-469,
477-483

Aspartic acid and asparagine hydroxylation site.

amino acids 262-274

FIGURE 37

CGGACGCGTGGGATTCAAGCAGTGGCCGTGGCTGCCAGAGCAGCTCCTCAGGGGAAACTAAG
CGTCGAGTCAGACGGCACCATATACTGCCTTAAAGTCGCTCGCCCTGCCGGCGCGTATC
CCCCGGCTACCTGGGCCGCCCCGGCGCGTGCAGCTGAGAGGGAGCGCGCGGGCAGCGA
GCCGCCGTGTGAGCCAGCGCTGCTGCCAGTGTGAGCGGGCTGTGAGGCCGGTGGGTGCGGA
GGGGCGTGTGCGGGCGCCGGCGCCCTGGGGTCAAACCCGAGCGTCTACGCTGCCATGA
GGGGCGCGAACCCCTGGGCCCACTCTGCCGTGCTGGCTGCCACCCAGCTCGCCGG
CAGCAGTCCCCAGAGAGACCTGTTTCACTGTGGTGGCATTCTACTGGAGAGTCTGGATT
TATGGCAGTGAGGTTTCTGGAGTGTACCCCTCCAAATAGCAAATGTACTTGGAAAATCA
CAGTCTCCGAAGAAAAGTAGCTGTTCTCAATTCCGATTATAGACCTCGAGAGTGACAAC
CTGTGCCGCTATGACTTGTGGATGTGTACAATGGCATGCCATGGCAGCGCATTGGCCG
CTTCTGTGCCACTTCCGGCTGGAGCCCTGTGTCAGTGGCAACAAGATGATGGTCAGA
TGATTTCTGATGCCAACACAGCTGGCAATGGCATGGCCATGTTCTCGCTGTAACCA
AACGAAAAGAGGGGATCACTATTGTGGAGGACTCTTGACAGACCTTCCGGCTCTTTAAAAC
CCCAACTGGCAGACCGGGATTACCCCTCAGGAGTCACCTGTGTGTGGCACATTGTAAGCC
CAAAGAATCAGCTTATAAGAATTTAGTTGAGAAGTTGATGTGGAGCGAGATAACTACTGC
CGATATGATTATGTGGCTGTGGTAAAGGGGGAGTCAACAGTGTAGAAGAAATTGGAAA
GTATTGTGGTGTAGTCACCTCGCCAATTGTGTCAGAGAAAATGAACCTTATTCACT
TTTATCAGACTTAAGTTAACTGAGGGTTATTGGTCACTACATATTCAAGGCAAAA
AAACTGCTTACAACACTACAGAACAGCGCTGCAACACCACATCCCTGTAACCACGGGTTAAA
ACCCACCGTGGCTTGTGTCACAAAAGGTAGACGGACGGGACTCTGGAGGGCAATTATT
GTTCAAGTGAATTGTATTAGCCGGACTGTTTACAAACCATCACTCGGATGGGAGTTG
CACGCCACAGTCGATCATCAACATCTACAAAGAGGGAAATTGGCGATTCAAGGCCGG
CAAGAACATGAGTCCCAGGTGACTGTGCTGCAAGCAGTGGCCCTCTCAGAACAGGTC
TAAATTACATTATTATGGGCCAAGTAGGTGAAGATGGGAGGGCAAATCATGCCAACAGC
TTAACATGATGTTCAAGGCAAGAACATGAGGCTCTGGATGCCCTTAAAATAAGCAATG
TTAACAGTGAACTGTGTCCATTAAAGCTGTATTCTGCCATTGCCATTGAAAGATCTATGTT
TCTCACTGAGAAAAAAATACTTATAAAATTACATATTCTGAAAGAGGATTCCGAAAGATGG
GACTGTTGACTCTTCACATGATGGAGGTATGAGGCCCTCCGAGATAGCTGAGGGAAAGTCTT
TGCTCTGCTGTGAGGGAGCAGCTATCTGATTTGGAAACACTGCCGACTTAACTGTGCGGTGATAGGA
AGCTAAAAGTGTCAAGCGTTGACAGCTTGAAGCGTTTATTATACATCTGTAAAGGAT
ATTAGAATTGAGTTGTGAGATGTCAAAAAAGATTAGAAGTGCATATTATAGT
GTTATTGTGTTTACCTTCAAGCCCTTGGCCCTGAGGTGTTACAAATCTTGTCTGCCCT
AACTCAATGCTTAATAAAATTTTAAAGGAAAAAA

FIGURE 38

MRGANAWAPLCLLLAATQLSRQQSPERPVFTCGGILTGESGFIGSEGFPGVYPPNSKCTWK
ITVPEGKVVVNLNFRFIDLESNDLCRYDFVDVYNGHANGQRIGRFCGTFRPGALVSSGNKMMV
QMISDANTAGNGFMAMFSAAEPNERGDQYCGGLLDRPSGSFKTPNWPDRDYPAGVTCVWHIV
APKNQLIELKFEKFDFVERDNYCRYDYVAVFNGGEVNDARRIGKYCGDSPAPIVSERNELLI
QFLSDLSLTADGFIGHYIFRPKLPTTTEQPVTTTFPVTTGLKPTVALCQQKCRRTGTLLEGN
YCSSDFVLAGTVITTRDGSLHATVSIINIYKEGNLAIQQAGKNMSARLTVVCKQCPLLRR
GLNYIIMGQVGEDGRGKIMPNSFIMMFKTKNQKLLDALKNKQC

Signal sequence:

amino acids 1-23

N-glycosylation site.

amino acids 355-359

Casein kinase II phosphorylation site.

amino acids 64-68, 142-146, 274-278

Tyrosine kinase phosphorylation site.

amino acids 199-208

N-myristoylation site.

amino acids 34-40, 35-41, 100-106, 113-119, 218-224, 289-295,
305-311, 309-315, 320-326, 330-336

Cell attachment sequence.

amino acids 149-152

FIGURE 39

CGGACGCGTGGCGGAGCGCTGGCGGCCACGGCGCCCGGGCTGGGGCGGTGCTTC
CCTTCTCCGTGGCTACGAGGGTCCCCAGCCTGGTAAAGATGGGCCCCATGGCCCCGAAGG
GCCTAGTCCAGCTGTCTGGGCTCAGCCTCTCCTCAACCTCCAGGACCTATCTG
CTCCAGCCCTCCACCTCCCCAGTCTTCTCCCCGCTCAGCCCCATCCGTGTCAACCTG
CCGGGGACTGGTTGACAGCTTAACAAGGGCTGGAGAGAACCATCCGGACAACTTGGAG
GTGAAACACTGCCCTGGGAGGAAGAGAATTGTCCAATACAAAGACAGTGAGACCCGCC
GTAGAGGTGCTGGAGGGTGTGTGCAAGTCAGACTTCGAGTGCACCCGCTGGAGCT
GAGTGAGGAGCTGGTGGAGAGCTGGTGGTTACAAGCAGCAGGAGGCCGACCTTCC
AGTGGCTGTGCTCAGATTCCCTGAAGCTCTGTCGCCCCGCAAGCACCTTCGGGCTCCTGC
CTTCCCCTGCTCTGGGGAACAGAGAGGCCCTGCGGTGGCTACGGGCAGTGTGAAGGAGAAGG
GACACGAGGGGGCACGGGCATGTGACTGCCAAGCCGGCTACGGGGTGAGGCCTGTGGCC
AGTGTGGCCTGGCTACTTGAGGCAGAACCAGCCAGGCCATCTGGTATGTTGGCTTGT
TTGGGCCCCTGTGCCGATGCTCAGGACCTGAGGAATCAAACTGTTGCAATGCAAGAAGG
CTGGGCCCCTGCATCACCTCAAGTGTGTAGACATTGATGAGTGTGGCACAGAGGGAGCCA
ACTGTGGAGCTGACCAATTCTGCGTAACACTGAGGGCTCCATGAGTGCGAGACTGTGCCAAG
GCCTGCCTAGGCTGCATGGGGCAGGGCCAGGTGCGTGAAGAAGTGTAGCCCTGGCTATCA
GCAGGTGGGCTCAAGTGTCTCGATGTGGATGAGTGTGAGACAGAGGTGTGCCGGAGAGA
ACAAGCAGTGTAAAACCGGGGCGGTATCGCTGCATGTGCCAGGGCTACAAGCAG
ATGGAAGGCATCTGTGTAAGGAGCAGATCCCAGAGTCAGCAGGCCTTCAGAGATGAC
AGAACAGCAGTTGGTGGTGTGCTGCAGCAGATGTTCTTGGCATCATCTGTGCACTGCCA
CGCTGGCTGCTAAGGGCACTGGTGTGTTACCGCCATCTTCATTGGGCTGTGCCGGCATG
ACTGGCTACTGGTGTGCAGAGCGCAGTGACCGTGTGCTGGAGGGCTCATCAAGGGCAGA
TATCGGGCCACCACCTGTTAGGACCTCCCTCCACCCAGCCTGCCAGAGCTGGGCTGCC
TCCTGCTGGACACTCAGGACAGCTGGTTATTTTGAGAGTGGGTAAGCACCCTACCTG
CCTTACAGAGCAGGCCAGGTACCCAGGCCGGCAGACAAGGCCCTGGGTAAAAGTAGC
CCTGAAGGTGGATACCAGAGCTCTCACCTGGCGGGACTGCCAGGCTCACAATGTTG
ATTTCAAAAGTTTTCTTAATGGTGGCTGCTAGAGCTTGGCCCTGCTTAGGATTAGGT
GTCCTCACAGGGTGGGGCCATCACAGCTCCCTCTGCCAGCTGCATGCTGCCAGTTCTGT
TCTGTGTTACCCACACCCATTGCCACTTATTATTCATCTCAGGAAATAAAGA
AAGGTCTGGAAAGTTAAAAAAAAAAAAAA

FIGURE 40

MAPWPPKGGLVPAVLWGLSLSFLNLPGPPIWLQPSPPPQSSPPPQPHPCHTCRGLVDSFNKGLER
TIRDNFGGGNNTAWEENNLSKYKDSETRLVEVLEGVCSKSDFECHRLLESELVESWWFHKQ
QEAPDLFQWLCSDSLKLCCPAGTFGPSCLPCCGGTERPCGGYGQCEGECTRGGSGHCDCQAG
YGEACGQCGCLGYFEAERNASHLVCSACFGPCARCSGPEESNCLQCKKGWALHHLKCVDIDE
CTGTGANCQADQFCVNTEGSYECRDCAKACLGCMGAGPGCRKKCSPGYQQVGSKCLDVDECE
TEVCPGENQCENTEGGYRCICAEGYKQMEGICVKEQIPESAGFFSEMTEDELVVLQQMFFG
III CALATLAAGDLDLVFTAIFIGAVAAMTGYWLSERSDRVLEGFIKGR

Signal sequence:

amino acids 1-29

Transmembrane domain:

amino acids 372-395

N-glycosylation site.

amino acids 79-83, 205-209

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 290-294

Casein kinase II phosphorylation site.

amino acids 63-67, 73-77, 99-103, 101-105, 222-226, 359-263

N-myristoylation site.

amino acids 8-14, 51-57, 59-65, 69-75, 70-76, 167-173, 173-179,
177-183, 188-194, 250-256, 253-259, 267-273, 280-286, 283-289,
326-332, 372-378, 395-401

Aspartic acid and asparagine hydroxylation site.

amino acids 321-333

EGF-like domain cysteine pattern signature.

amino acids 181-193

FIGURE 41

TGAGACCCCTCTGCAGCCTTCAGGGACAGCCCCACTCTGCCCTTGCTCCTCCAGGGCA
GCACC**ATG**CAGCCCCCTGTGGCTCTGCTGGGACTCTGGGTGTTGCCCTGGCAGCCCCGGG
GCCGCCCTGACGGGGAGCAGCTCCTGGCAGCCTGCTGCCAGCTGAGCTCAAAGAGGT
GCCACCCCTGGACAGGGCGACATGGAGGAGCTGGTCACTCCCACCCACGTGAGGGCCAGT
ACGTGGCCCTGCTGCAGCGCAGCCACGGGACCGCTCCCGCAAAGAGGTTCAAGCCAGAGC
TTCCGAGAGGTGGCCGGAGGTTCTGGCGTTGGAGGCCAGCACACACCTGCTGGTTCGG
CATGGAGCAGCGGCTGCCGCCAACAGCGAGCTGGTGCAGGCCGTGCTGCCGTCTTCAGG
AGCCGGTCCCCAAGGCCGCGCTGCACAGGCACGGCGGCTGTCAGGCCGAGGCCGGGC
CGGGTACCGCTGAGTGGCTGCCGCTCCGCACAGGCTCAACCGCACCTCCCTCATCGA
CTCCAGGCTGGTGTCCGTCACGAGCGGCTGGAAGGCCCTGACGTGACCGAGGCCGTGA
ACTTCTGGCAGCAGCTGAGCCGCCCGCACCGCCTGCTGTCACAGGTGTCGGTGCAGAGG
GAGCATCTGGCCCGCTGGCGTCCGGGCCACAAGCTGGTCCCTTGCCCTGCCAGGGGGC
GCCAGCCGGCTGGGAGCCCCAGCTGGAGCTGCACACCCGGACCTGGGACTATGGAG
CTCAGGGCAGTGTGACCCCTGAAGCACCAATGACCGAGGGCACCCGCTGTCGCCAGGAG
ATGTACATTGACTCGCAGGGGATGAAGTGGGCCAGAATGGGTGCTGGAGCCCCGGCTT
CCTGGCTTATGAGTGTGTCGGCACCTGCCGAGCCCCGGAGGCCCTGCCCTCAAGTGGC
CGTTCTGGGGCTCGACAGTGATCGCCTGGAGACTGACTCGCTGCCATGATCGTCAGC
ATCAAGGAGGGAGGCAGGACCAGGCCAGTGGTCAGCTGCCAACATGAGGGTGCAGAA
GTGAGCTGTGCCCTGGATGGTGCCTCGGCCAAGGAGGCCAT**AGGCGCTAGTG**
TAGGCCATCGAGGGACTTGAATTGTGTTCTGAAGTGTGAGGGTACCAAGGAGAGCTG
GCGATGACTGAACGTGCTGATGGACAAATGCTCTGCTCTAGTGAGGCCCTGAATTGCTT
CCTCTGACAAGTTACCTCACCTAATTTTGCTTCTCAGGAATGAGAAATCTTGCCACTGGA
GAGGCCCTGCTCAGTTCTCTATTCTTATTCACTGCACTATATTCTAACGACTTACAT
GTGGAGATACTGTAACCTGAGGGCAGAAAGCCANTGTGTCATTGTTACTTGCTCTGTCAC
TGGATCTGGCTAAAGTCTCCACCACCACTGGACCTAACGACCTGGGTTAAGTGTGGGT
TGTGATCCCCAATCCAGATAATAAGACTTGTAAAACATGAATAAAACACATTATTCT
AAAA

FIGURE 42

MQPLWLCWALWVLPLASPGAAALTGEQLLGSSLRQLQLKEVPTLDRADMEELVIPTHVRAQYV
ALLQRSHGDRSRGKRFQSFSREVAGRFLALEASTHLLVFGMEQRLPPNSELVQAVLRFQEP
VPKAALHRHGRLSPRSARARVTVEWLVRDDGSNRTSLIDSRLVSVHESGWKAFCVDTEAVNF
WQQLSRPRQPPLLQVSQREHLGPLASGAHKLVRFASQGAPAGLGEPEPQLEHTLDLGDYGAQ
GDCDPEAPMTEGTRCCRQEMYIDLQGMKWAENWVLEPPGFAYECVGTCRQPPEALAFKWPF
LGPRQCIASETDSLPMIVSIKEGGRTRPQVVSLPNMRVQKCSCASDGALVPRRLQP

Signal sequence:

amino acids 1-18

N-glycosylation site.

amino acids 158-162

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 76-80

Casein kinase II phosphorylation site.

amino acids 68-72, 81-85, 161-165, 169-173, 319-323, 329-333

N-myristoylation site.

amino acids 19-25, 156-162, 225-231, 260-266, 274-280

Amidation site.

amino acids 74-78

TGF-beta family signature.

amino acids 282-298

FIGURE 43

TRANSCRIPTION

GTCTGTTCCCAGGAGTCCTCGGGCTGTTGTCAGTGGCCTGATCGCGAGGGGACAAA
GGCGCAAGTCGAGAGGAAACTGTTGTCCTCATATTGGCGATCCTGTTGCTCCCTGG
CATTGGGCAGTGTACAGTCACCTCTGAACCTGAAGTCAGAATTCTGAGAATAATCCT
GTGAAGTTGTCCTGTCCTACTCGGGCTTTCTCTCCCGTGGAGTGGAAAGTTGACCA
AGGAGACACCACCAAGACTCGTTGCTATAATAACAAGATCACAGCTTCTATGAGGACCGGG
TGACCTTCTGCCAAGTGGTACACCTTCAGTCCGTCAGACACGGGAAGACACTGGGACATAC
ACTTGTATGGTCTCTGAGGAAGGGCGCAACAGCTATGGGGAGGTCAGGTCAAGGCTCATCGT
GCTTGTGCCCTCATCCAAGCCTACAGTTAACATCCCTCCTGCCCCACATTGGGAACCGGG
CACTGCTGACATGCTCAGAACAGATGGTCCCCACCTTCTGAATAACACCTGGTTCAAAGAT
GGGATAGTGTGCCCTACGAATCCAAAAGCACCCGTGCCTTCAGCAACTCTCCTATGTCCT
GAATCCCACACAGGAGAGCTGGCTTTGATCCCTGTCAGCTCTGATAACTGGAGAATACA
GCTGTGAGGCACCGGAATGGGTATGGGACACCCATGACTTCAAATGCTGCGCATGGAAAGCT
GTGGAGCGGAATGTGGGGTCTCGTGGCAGCGTCCTGTAACCTGATTCTCCTGGGAAT
CTTGGTTTTGGCATCTGGTTGCCATAGCCGAGGCCACTTGTACAGAACAAAGAAAGGG
CTTCGAGTAAGAAGGTGATTACGCCAGCCTAGTGCCTGAAGTGAAGGAGATTCAAACAG
ACCTCGTCATTCTGGTGTGAGCTGGTCGGCTCACCGCCTATCATCTGCAATTGCGCTTACT
CAGGTGCTACCGGACTCTGGCCCTGATGTCGTAGTTCACAGGATGCCTTATTGTCTC
TACACCCACAGGGCCCCCTACTTCTCGGATGTGTTTAATAATGTCAGCTATGTCCCC
ATCCTCCCTCATGCCCTCCCTCCCTACCACTGCTGAGTGGCCTGGAACATTGTTAAA
GTGTTTATTCCCCATTCTTGGAGGGATCAGGAAGGAATCTGGGTATGCCATTGACTTCCC
TTCTAAGTAGACAGCAAAATGGGGGGTCGAGGAATCTGCACTCAACTGCCACCTGGC
TGGCAGGGATCTTGAAATAGGTATCTTGAGCTTGGTCTGGCTCTTCTGTGTACTGAC
GACCAAGGGCCAGTGTCTAGAGCGGGATTAGAGGCTAGAGCGGCTGAAATGGTTGG
TGATGACACTGGGTCTTCCATCTCTGGGCCACTCTCTGTCTTCCATGGGAAGTG
CCACTGGGATCCCTCTGCCCTGCTCTGAAATACAAGCTGACTGACATTGACTGTCTGT
GGAAAATGGGAGCTTGTGAGGAGCATAGTAATTTCAAGAGAACCTGAAGCCAAAAG
GATTTAAAACCGCTCTGCTCTAAAGAAAAGAAAATGGAGGCTGGCGCAGTGGCTCACGCC
TAATCCCAGAGGCTGAGGCAGCGGATCACCTGAGGTGGAGGATCAGCCTGACCA
ACATGGAGAACCCACTGGAAATACAAGTTAGCCAGGCATGGTGGTGCATGCCGTAGTC
CCAGCTGCTCAGGAGCCTGGCAACAGAGCAAAACTCCAGCTCAAAAAAA

FIGURE 44

MGTKAQVERKLLCLFILAILLCSLALGSVTVHSSEPEVRIPENNPVKLSCAYSGFSSPRVEW
KFDQGDTTRLVCYNNNKITASYEDRVTFLPTGITFKSVTREDTGTYTCMVSEEGGN SYGEVKV
KLIVLVPPSKPTVNI PSSATI GNR A VLT C SE QDG S P P S E Y T W F K D G I V M P T N P K S T R A F S N S
SYVLNPPTT GELVFDPLSASDTGEYSCEARNGYGT PMTSNAVRMEAVRN VGV VIVA AVL VTLI
LLGILVFGIWFAYSRGHFDRTKKGTSSKKVIYSQPSARSEGFKQTSSFLV

Signal sequence:

amino acids 1-27

Transmembrane domain:

amino acids 238-255

N-glycosylation site.

amino acids 185-189

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 270-274

Casein kinase II phosphorylation site.

amino acids 34-38, 82-86, 100-104, 118-122, 152-156, 154-158,
193-197, 203-207, 287-291

N-myristoylation site.

amino acids 105-111, 116-122, 158-164, 219-225, 237-243, 256-262

FIGURE 45

CAGCGCGTGGCCGGCGCGTGTGGGACAGCATGAGCGCGGTTGGATGGCGCAGGTTGGA
GGCTGGCGAACAGGGCTCTGGCCTGGCGCTGCTGCTGCTCGGCCTCGGACTAGGCCT
GGAGGCCGCGAGGCCGTTCCACCCCGACCTCTGCCAGGCCAGGCCAGCTCG
GCTGTGCCACCACCAAGTCCAGTGCGCACCGAGGGCTATGCGTGCCCCCTCACCTGG
CGCTGCGACAGGGACTTGGACTGCAGCGATGGCAGCGATGAGGAGGAGTCAGGATTGAGCC
ATGTACCCAGAAAGGGCAATGCCACGCCCTGGCCTCCCTGCCCTGCACCGCGTCA
GTGACTGTCTGGGGAACTGACAAGAAACTGCGCAACTGCAGGCCCTGGCCTGCCTAGCA
GGCGAGCTCCGTTGCACGCTGAGCGATGACTGCATTCCACTCACGTGGCGCTGCGACGCCA
CCCAGACTGTCCGACTCCAGCGACGAGCTGGCTGTGAGAACATGAGATCCTCCCGAAG
GGGATGCCAACCATGGGCCCCCTGTGACCCCTGGAGAGTGTACCTCTCTCAGGAATGCC
ACAACCATGGGCCCCCTGTGACCCCTGGAGAGTGTCCCTCTGTGGAAATGCCACATCCTC
CTCTGCCGAGACCAGTCTGGAAGCCAACGCCTATGGGTTATTGCAGCTGCTGCCGTGC
TCAGTGAAGCCTGGCACCGGACCCCTCCCTTTGTCTGGCTCCGAGCCAGGAGCGC
CTCCGCCACTGGGTTACTGGTGGCCATGAAGGAGTCCCTGCTGCTGTAGAACAGAACAG
CTCGCTGCCCTGAGGACAAGCACTTGCCACCCGTCACTCAGGCCCTGGCGTAGCCGGACA
GGAGGAGAGCAGTGTGCGGATGGTACCCGGCACACCGCCCTCAGAGACCTGAGTTCTT
CTGGCCACGTGGAACCTCGAACCCGAGCTCTGCAGAACAGTGGCCCTGGAGATTGAGGGTCCC
TGGACACTCCCTATGGAGATCCGGGAGCTAGGATGGGAAACCTGCCACAGCCAGAACAG
GGGCTGGCCCCAGGCAGCTCCAGGGTAGAACGCCCTGTGCTTAAGAACACTCCCTGCTG
CCCCGTCTGAGGGTGGCGATTAAAGTTGCTTC

FIGURE 46

MSGGWMAQVGAWRTGALGLLLLLLGLGLLEAAASPLSTPTSAQAAGPSSGSCPPTKFQCR
TSGLCVPLTWRCRDLDCSDGSDEEECRIEPCTQKGQCPPPGLPCPCTGVSDCSGGTDKKL
RNCSRLAACLAGELRCTLSDDCIPLTWRCGDGHPDPCPDSSDELGCGTNEILPEGDATTMGPV
LESVTSLRNATTMGPVTLIESVPSVGNATSSAGDQSGSPTAYGVIAAAAVLSASLVTATLL
LLSWLRAQERLRLPLGLLVAMKESLLLSEQKTSLP

Signal sequence:

amino acids 1-30

Transmembrane domain:

amino acids 230-246

N-glycosylation site.

amino acids 126-130, 195-199, 213-217

Casein kinase II phosphorylation site.

amino acids 84-88, 140-144, 161-165, 218-222

N-myristoylation site.

amino acids 3-9, 10-16, 26-32, 30-36, 112-118, 166-172, 212-218,
224-230, 230-236, 263-269

Prokaryotic membrane lipoprotein lipid attachment site.

amino acids 44-55

Leucine zipper pattern.

amino acids 17-39

FIGURE 47

CCACCGTCCGGCTCGCTCGCTCGCAGCGCGGAGGAGGAGGAAGGAAAGCTGCATGCATGAGACCCACAGA
CTCTTGCAAGCTGGATGCCCTGTGGATGAAAGAATGTATCATGGAATGAACCGAGCAATG
GAGATGGATTCTAGAGCAGCAGCAGCAGCACACCTCAGTCCCCAGAGACTCTTG
GCCGTGATCCTGTGGTTCACTGGCCTGTGCTTCGGCCCTGCACAGCTCACGGCGGGT
CGATGACCTTCAAGTGTGCTGACCCCGCATTCCCGAGAAATGGCTCAGGACCCAGCG
GAGGGGTTCTTGAAAGGCTGTAGCCGATTTCACTGCCAAGACGGATTCAAGCTGAAG
GGCCTACAAAGAGACTGTGTTGAAGCATTAAATGAAACCTAGGCTGGATCCAAGTGA
TAATTCCATCTGTGTGCAAGAAGATTGGCGTATCCCAAATCGAAGATGCTGAGATTATA
ACAAGACATATAGACATGGAGAGAACGCTAACATCATCACTGTGATGAAGGATTCAAGATCCGG
TACCCCCACCTACAAATATGGTTCTATTATGTCGGATGATGGAACGTGGAATAATCTGCC
CATCTGCAAGGCTGCCTGAGACCTCTAGCCTTCTAATGGCTATGTAACATCTGAGC
TCCAGACCTCCTCCGGTGGGACTGTGATCCTATCGCTGCTTCCGGATTTAACCT
GATGGGCTGCGTATCTGAGTGCTTACAAAACCTTATCGGTGTCAGCCCACCCGGTG
CCTGCTCTGGAAGCCAAGTGTCCACTACCTCAAATGGTAGTCACGGAGATTTCTGCT
GCCACCCGCGGCCCTGTGAGCGTACAACCACCGAACGTGGTGGAGTTTACTGCGATCCT
GGCTACAGCCTACCAGCGACTACAAGTACATCACCTGCCAGTATGGAGAGTGGTTCCCTC
TTATCAAGTCTACTGCATCAAATCAGAGCAAACGTGGCCAGCACCCATGAGACCCCTCTGA
CCACGTGGAAGATTGTGGCGTTCACGGCAACCAGTGTGCTGGTGCTGCTCGTCATC
CTGGCCAGGATGTCCAGACCAAGTCAAGGCCACTTCCCCCAGGGGCTCCCGAG
TTCCAGCAGTGAACCTGACTTGTGGTGGTAGACGGCGTGGCGTCACTGCTCCGTCCTATG
ACGAAGCTGTGAGTGGCGGTTAGTGCCTTACGGCCGGTACATGGCCTCTGTGGCCAG
GGCTGCCCTTACCGTGGACGACCAGAGCCCCCAGCATAACCCGGCTCAGGGACACGGA
CACAGGCCAGGGAGTCAGAAACCTGTGACAGCGTCTCAGGCTCTGAGCTGCTCCAAA
GTCTGTATTCAACCTCCAGGTGCAAGAGAGCACCCACCCCTGCTCGGACAACCTGACATA
ATTGCCAGCACGGCAGAGGAGGTGGCATCCACCAAGCCAGGCATCCATCATGCCCACTGGT
GTTGTTCTAAGAAACTGATTGATTAAAAAATTCCCAAAGTGTCTGAAGTGTCTTCAA
ATACATGTTGATCTGTGGAGTTGATTCTTCTCTTGGTTTAGACAATGTAACAA
AGCTCTGATCTTAAATTGCTATGCTGATAGAGTGGTGAGGGCTGGAAGCTTGATCAAGTC
CTGTTCTTCTTGACACAGACTGATTAAAAATTAAAAGNAAAAAA

FIGURE 48

MYHGMNPSNGDFLEQQQQQQQQPQPQRLLAVILWFQLALCFGPAQLTGGFDDLQVCADPGI
PENGFRTPSGGVFFEGSVARFHQCQDGFKLKAGATKRLCLKHFNGLTGWI PSDKNSICVQEDCRI
PQIEDAEIHNKTYRHGEKLIITCHEGFKIRYPDLHNMSLCRDDGTWNNLPICGCLRPLAS
SNGYVNISELQTSPVGTVISYRCFPGFKLDSAYLECLQNLIWSSSPRCLALEAQVCPLP
PMVSHGDFVCHPRPCERYNHGTVEFYCDPGYSLTSDYKYITCQYGEWFPSYQVYCIKSEQT
WPSTHETLLTTWKIVAFTATSVLLVLLVILARMFQTKFKAHFPPRGPPRSSSDPDFVVVD
GVPVMLPSYDEAVSGGLSALGPGYMASVGQGCPLPVDDQSPPAYPGSGD TDTGPGESETCDS
VSGSSELLQSLYSPPRCQESTHPASDNPDIIASTAEEVASTSPGIHHAHWVLFLRN

Signal sequence:

amino acids 1-41

Transmembrane domain:

amino acids 325-344

N-glycosylation site.

amino acids 104-108, 134-138, 192-196

Casein kinase II phosphorylation site.

amino acids 8-12, 146-150, 252-256, 270-274, 313-317, 362-366,
364-368, 380-384, 467-471, 468-472

N-myristoylation site.

amino acids 4-10, 61-67, 169-175, 203-209, 387-393, 418-424,
478-484

Prokaryotic membrane lipoprotein lipid attachment site.

amino acids 394-405

FIGURE 49

CCCCACGCGTCCGCTCCGC~~GCCCCTCCCCCCC~~GCCTCCCGTGC~~GGTCCGTGGTGGCCTAGAGA~~
TGCTGCTGCCGCCGGTGCAGTTGTCGCGCACGCC~~T~~C~~T~~G~~CCCGCCAGCCCGCTCCACC~~CGCT
AGCGCCCGAGTGT~~CGGGGGCG~~ACCCGAGT~~GGG~~~~A~~~~T~~~~GAGGCCGGAAACCGCGCTACAGG~~
CCGTGCTGCCGCCGGTGC~~TGGTGGGCTGCGGCCGCGACGGGTCGCTGAGTGCC~~
TCGGATTTGGACCTCAGAGGAGGGCAGCCAGT~~C~~TG~~CCGGGAGGGACACAGAGCCTTGT~~TA
TAAAGTCATTACTTCCATGATACTTCTGAAGACTGA~~ACTTTGAGGAAGCCAAGAACAGCCT~~
GCAGGGAGGGATGGAGGCCAGTAGTCAGCATCGAGT~~C~~TGAAGATGAACAGAA~~ACTGATAGAA~~
AAGTT~~CATTGAAAACCTT~~GGCCATCTGATGGT~~GACTTCTGGATTGGCTCAGGAGGCGTGA~~
GGAGAAACAAAGCA~~ATAGC~~ACGCC~~TGCCAGGAC~~CTTATGCTTGA~~CTGACTGATGGCAGCATAT~~
CACAATTTAGGA~~ACTGGTATG~~GGATGAGGCC~~TCTGCGGCAGCGAGGTCTGCGTGGT~~CATG
TACCATCAGCC~~CATCGG~~CACCCGCTGGCATGGAGGCC~~CTACATGTT~~CAGT~~GAGAAACAGCAGTT~~CTT
CCGGT~~GCAACATGAAGAACAA~~TTTCAT~~TGCAAATATTCTGATGAGAAACACAGCAGTT~~CTT
CTAGAGAAGCTGAAGGT~~GAGGAAACAGAGCTGACAA~~CACCTGTACTTCCAGAAGAA~~ACACAG~~
GAAGAAGATGCC~~AAAAAAACATT~~AAAGAAAGTAGAGAAGCTG~~CC~~TTGAATCTGGC~~CTACAT~~
CTTAATCCCAG~~CATT~~CCC~~TCTCCTCC~~CTTGTGGT~~CACCACAGTTGATGTTGGTTT~~
GGATCTGTAGAAAAGAAAAGGGAGCAGCCAGAC~~CTAGC~~ACAAAGAAGCAACACACC~~ATC~~
TGGCC~~CTCTC~~ACCAGGGAAACAGCCGGAC~~TAGAGG~~T~~C~~AC~~ATGT~~C~~ATAAGAAACAA~~
AA~~AGCGAAGCTGACTT~~AGCTGAGACCCGGCAGAC~~CTGAA~~GAAGA~~ATATT~~T~~CCGAGTGT~~TT
CGGGAGAAGCC~~ACT~~CCCGATGAC~~ATGT~~CTTGT~~GACTATGACAACATGGCTGT~~GAACCC~~ATCA~~
GAAAGTGGGTTTG~~GACTCTGGT~~GAGCGT~~GGAGGTGGATT~~T~~GACCAATGACATT~~T~~ATG~~
GTTCTCCCAGAC~~AAATGGGGAGGAGT~~AAAGGAGT~~CTGGATGGGTGGAA~~ATGAA~~ATATATG~~
GTT~~TAGGACATATA~~AAA~~ACTGAACTGACAACAA~~T~~GGAAAAGAAATGATAAGC~~AA~~ATC~~
CTCTT~~ATT~~TTCTATAAGGAAA~~ACACAGAAGGT~~T~~CTATGAAACAAGCTT~~AGATCAGGT~~CTGT~~
GGATGAGC~~ATGT~~GGT~~CCCCACGAC~~CTCTG~~TTGGAC~~CCCCACG~~TTGGCTGT~~T~~ATC~~CTT~~AT~~
CCCAGCGAGT~~CATCC~~AGCTG~~ACCTT~~T~~ATGAGAAGGT~~AC~~CTTGCCAGGT~~CTGGC~~CACATAG~~
GAGT~~CTCAAA~~AT~~GTCACTTGGTTGT~~T~~ATCTAAC~~TTAAGGGACAGAGC~~GT~~TT~~ACCTG~~
GCAGTGATAAAAGATGGG~~CTGTGGAGCTTGGAAAACCAC~~C~~CTGT~~TT~~CC~~TTG~~CT~~T~~ATACAG~~
CAGCACAT~~ATTAT~~C~~ATACAGACAGAAAATCCAGA~~AT~~CTTTCAAAGCCCACATATGGTAGCAG~~
GTTGGC~~CTGTG~~C~~ATCGG~~AA~~TTCTCATATCTGTTTTT~~CAAAGA~~ATAAAATCAAATAAAGA~~
GCAGG~~AAAAAA~~

FIGURE 50

MRPGTALQAVLLAVLLVGLRAATGRLLSASDDLRRGGQPVCRGGTQRPCYKVIYFHDTSRRL
NFEEAKEACRRDGGQLVSIESEDEQKLIEKFIEENLLPSDGFWIGLRRREEKQSNSTACQDL
YAWTDGSISQFRNWYVDEPSCGSEVCVVMYHQPSAPAGIGGPYMFQWNDRNCNMKNNFICKY
SDEKPAVPSREAEGEETELTPVLPEETQEEDAKTFKESREAALNLAYILIPSIPLLLLLV
VTTVVCWVWICRKRKREQPDSTKKQHTIWPSPHQGNSPDLEVNVIRKQSEADLAETRPDL
KNISFRVCSCGEATPDDMSCDYDNMAVNPSSESGFVTLSVSEGFVTNDIYEFSPDQMGRSKES
GWVENEIYGY

Signal sequence:

amino acids 1-21

Transmembrane domain:

amino acids 235-254

N-glycosylation site.

amino acids 117-121, 312-316

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 296-300

Casein kinase II phosphorylation site.

amino acids 28-32, 30-34, 83-87, 100-104, 214-218, 222-226,
299-303, 306-310, 323-327

N-myristoylation site.

amino acids 18-24, 37-43, 76-82, 146-152

FIGURE 51

GGGGTCTCCCTAGGGCGGGAGGCACAGCGGTCCCTGCTGAAGGGCTGGATGTACGC
ATCCGCAGGTTCCCGCGACTTGGGGCGCCCGCTGAGCCCCGGCGCCCGAGAAGACTTGT
GTTTGCCTCCTGCAGCCTCAACCCGGAGGGCAGCGAGGGCTACCACCATGATCATGGTGT
GTTCA~~G~~CAGCATGCGCTTGTGGACCCCAGTGGCGTCTGACCTCGCTGGCGTACTGCCTGCAC
AGCGGGCGGGTGGGCCCTGGCCGAGCTGCAGGAGGCCAGTGGCGTACTGTCCGGTCGACCGCAGC
CTGCTGAAGTTGAAATGGTGAGGTGCTGTTGACACGGGGCTGGAGTCTCTCAAGCC
GCTCCCCTGGAGGGCAGGTAGAGTGGAACCCCCAGCTATTAGAGGTCCACCCCCAACTC
AGTTTGAATTACACAGTCACCAATCTAGCTGGTGGTCCGAAACCATATTCTCCTACGACTCT
CAATACCATGAGACCACCCCTGAAGGGGGCATGTTGCTGGCAGCTGACCAAGTGGCAT
GCAGCAATGTTGCTGGAGAGAGACTGAGGAAGAACTATGTGGAGACATTCCCTTC
TTTCAACCAACCTCAACCCACAGGAGTCTTATTCTGTTCACTAACATTTTGGAACTCTG
GAGTCCACCCGTTGTTGCTGGCTGGCTTTCCAGTGTAGAAAGAAGGACCCATCATCAT
CCACACTGATGAAGCAGATTCAAAGTCTGTATCCAAACTACCAAAGCTGCTGGAGGCTGA
GGCAGAGAACCGAGGGCCGGAGGCAGACTGCCTTTACAGCCAGGAATCTCAGAGGATTG
AAAAAGGTGAAGGACAGGATGGCATTGACAGTAGTGTATAAGTGGACTTCTCATCCTCCT
GGACAACGTTGGCTGCCGAGCAGGCACACAACTCCAAAGCTGCCCCATGCTGAAGAGATTG
CACGGATGATCGAACAGAGAGCTGTGGACACATCCTGTACATACTGCCAAGGAAGACAGG
GAAAGTCTCAGATGGCAGTAGGCCATTCCACATCCTAGAGAGCAACCTGCTGAAAGC
CATGGACTCTGCCACTGCCCGACAAGATCAGAAAGCTGTATCTCATGCCGCTATGATG
TGACCTTCATACCGCTCTTAATGACCTGGGATTTTGACCAAAATGCCACCGTTGCT
GTTGACCTGACCATGGAACCTTACCAAGCACCTGGAACTCTAAGGAGTGTTGTGCAGCTCTA
TTACACGGGAAGGGAGCAGGTGCCAGGGTGGCAGAGGTGCTGATGGCTCTGCCGCTGGACATGT
TCTTGAATGCCATGTCAGTTATACCTTAAGCCCAGAAAATACCATGCACTCTGCTCTCAA
ACTCAGGTGATGGAAGTTGGAAATGAAGAGTACTGATTATAAAAGCAGGATGTGTTGATT
TTAAAAATAAGTGCCTTATACAATG

FIGURE 52

MITGVFSMRLWTPGVVLTSLAYCLHQRRVALAELQEADGQCPVDRSLLKLKMVQVVFRHGR
SPLKPLPLEEQQVEWNPQLLEVPPQTQFDYTVNLAGGPKFYSFYDSQYHETTLKGGMFAGQL
TKVGMQQMFAFLGERLRKNYVEDIPFLSPTFNPQEVFIRSTNIFRNLESTRCLLQAGLFQCQKE
GPIIHTDEADSEVLYPNYQSCWSLRQRTRGRRQTASLQPGISEDLKKVKDRMGIDSSDKVD
FFILLDNVAEEQAHNLPCPMLKRFARMIEQRAVDTSLYILPKEDRESLQMAVGFLHILES
NLLKAMDSATAPDKIRKLYLYAAHDVTFIPLLMTLGIFDHKWPPFAVDLTMELYQHLESKEW
FVQLYYHGKEQVPRGCPDGLCPLDMFLNAMSVTLSPEKYHALCSQTQVMEVGNEE

Signal sequence:

amino acids 1-23

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 218-222

Casein kinase II phosphorylation site.

amino acids 87-91, 104-108, 320-324

Tyrosine kinase phosphorylation site.

amino acids 280-288

N-myristoylation site.

amino acids 15-21, 117-123, 118-124, 179-185, 240-246, 387-393

Amidation site.

amino acids 216-220

Leucine zipper pattern.

amino acids 10-32

Histidine acid phosphatases phosphohistidine signature.

amino acids 50-65

FIGURE 53

CTCCTCTAACATACTTGAGCTAAAACATAAATATTGCTGCTGGGGACCTCCTCTAGCCT
TAAATTCAGCTCATCACCTCACCTGCCTGGTCATGGCTCTGCTATTCTCCTTGATCCTT
GCCATTGACCAGACCTGGATTCCTAGCGTCTCCATCTGGAGTGCAGCTGGTGGGGGCCT
CCACCGCTGTGAAGGGCGGGTGGAGGTGGAACAGAAAGGCAGTGGGGCACCGTGTGATG
ACGGCTGGGACATTAAAGGACGTGGCTGTGTTGTGCCGGAGCTGGCTGTGGAGCTGCCAGC
GGAACCCCTAGTGGTATTTGTATGAGCCACAGCAGAAAAAGAGCAAAAGGTCTCATCCA
ATCAGTCAGTGCACAGGAACAGAAGATACTGGCTCAGTGTGAGCAAGAAGAAGTTATG
ATTGTTACATGATGAAGATGCTGGGGCATCGTGTGAGAACCCAGAGAGCTTTCTCCCCA
GTCCCAGAGGGTGTCAAGGCTGGCTGACGCCCTGGCATTCGAAGGGACCGTGGAAAGTGAA
GCACCAAGAACAGTGGTATACCGTGTGCCAGACAGGCTGGAGCCTCCGGCCGCAAAGTGG
TGTGCCGGCAGCTGGATGTGGAGGGCTGTACTGACTCAAAACGCTGCAACAAGCATGCC
TATGGCGAAACCCATCTGGCTGAGCCAGATGTCACTGCTCAGGACGAGAAGCAACCCCTCA
GGATTGCCCTCTGGCCTTGGGAGAACACCTGCAACCATGATGAAGACACGTGGTGC
AATGTGAAGATCCCTTGACTTGAGACTAGTAGGAGGAGACAAACCTCTGCTCTGGCGACTG
GAGGTGCTGCACAAGGGCTATGGGCTCTGCTGTGATGACAACGGGAGAAAAGGAGA
CCAGGTGGTATGCAAGCAACTGGCTGTGGGAAGTCCCTCTCCCTCAGAGACCGGA
AATGCTATGCCCTGGGTTGGCCGCATCTGGCTGGATAATGTTGCTGCTCAGGGAGGAG
CAGTCCTGGAGCAGTGCAGCACAGATTGGGTTCAAGACTGCACCCACAGGAAGA
TGTGGCTGTCACTGCTCAGTGTAGGTGGCATCATCTAATCTGTTGAGTGCCTGAATAGAA
GAAAAACACAGAAGAAGGGAGCATTTACTGCTCATGACTGCATGGGATGAACACTGATCT
TCTTCTGCCCTGGACTGGGACTTATACCTGGTGCCTGATTCTCAGGCCCTCAGAGTGG
ATCAGAACTTACAACATCAGGTCTAGTCTCAGGCCATCAGACATAGTTGGAACATAC
CCACCTTCTATGTCCTCACATTGCACACAGCAGATTCCAGCCTCCATAATTGTTGTT
CAACTACTAAATACATTCTCACACACACACACACACACACACACACACACACACATA
CACCAATTGTCCTGTTCTGAAAGAACTCTGACAAAATACAGATTGGTACTGAAAGAGA
TTCTAGAGGAACGGAAATTAAAGGATAATTCTGAATTGTTATGGGTTCTGAAATTG
GCTCTATAATCTAATTAGATATAAAATTCTGTAACCTTATTACAATAAAAGATAGCAC
TATGTTCAAA

FIGURE 54

MALLFSLILAICTRPGFLASPSGVRLVGGHLRCEGRVEVEQKGQWGTVCDDGWDIKDVAVLC
RELGCGAASGTPSGILYEPPAEKEQKVLIQSVSCTGTEDTLAQCEQEEVYDCSHDEDAGASC
ENPESSFSPVPEGVRLADGPGHCKGRVEVKHQNQWYTVCQTGWSLRAAKVVCRQLGCGRAVL
TQKRCNKHAYGRKPIWLSQMCSGREATLQDCPSGPWGKNTCNHDEDTWVECEDPFDLRLVG
GDNLCGSRLEVHKGVWGSVCDDNWGEKEDQVVCKQLGCGKSLSPSFRDRKCYGPGVGRIGL
DNVRCSGEEQSLEQCQHHRFWGFHDCTHQEDVAVICSV

Signal sequence:

amino acids 1-15

Casein kinase II phosphorylation site.

amino acids 47-51, 97-101, 115-119, 209-213, 214-218, 234-238,
267-271, 294-298, 316-320, 336-340

N-myristoylation site.

amino acids 29-35, 43-49, 66-72, 68-74, 72-78, 98-104, 137-143,
180-186, 263-269, 286-292

Amidation site.

amino acids 196-200

Speract receptor repeated domain signature.

amino acids 29-67, 249-287

FIGURE 55

ACTGCACTCGGTTCTATCGATTGAATTCCCCGGGGATCCTCTAGAGATCCCTGACCTCGAC
CCACCGCGTCCGGGACCGTGGCGGACCGCTGGGCGGCCTACCGAGAAGAGTCGCGGAAG
GTGAAGGCCATGGACTTCATCACCTCCACAGCCATCCTGCCCTGCTGTTGGCTGCCCTGGG
CGCTTCTCGGCTCTTCCGGCTGCAGTGGGTGCGCGGGAAAGGCCACTCGGGAATGCTG
TGGTGGTGTACAGGGGCCACCTCAGGGCTGGGCAAAGAAATGTGCAAAGAGCTTCTATGCT
GCGGGTGCTAAACTGGTGTCTGTGGCCGAATGGTGGGGCCCTAGAAGAGCTCATCAGAGA
ACTTACCGCTTCTCATGCCACCAAGGTGCAGACACACAAGCCTTACGGTGGACCTTCGACC
TCACAGACTCTGGGCCATAGTTGCAAGCAGCTGAGATCCTGCAGTGCTTGGCTATGTC
GACATACTTGTCAACAATGCTGGGATCAGCTACCGTGGTACCATCATGGACACCAAGTGG
TGTGGACAAGAGGGTCAAGGAGAACAAACTACTTGGCCAGTTGCTCTAACGAAAGCACTCC
TGCCTCCATGATCAAGAGGGCAAGGCCACATTGTCGCCATCAGCAGCATCCAGGGCAAG
ATGAGCATTCTTTGATCAGCATATGCAAGCCTCCAAGCACGCAACCCAGGCTTCTTGA
CTGCTGCGTGCCAGATGGAACAGTATGAAATTGAGGTGACCGTCATCAGCCCCGGCTACA
TCCACACCAACCTCTGTAAATGCCATCACCGGGATGGATCTAGGTATGGAGTTATGGAC
ACCACACAGCCAGGGCGAACCCCTGTGGAGGTGGCCAGGATGTTCTGCTGCTGG
GAAGAAGAAGAAAATGTGATCTGGCTGACTTAAGCTGCTTCTGGCTGTTATCTCGAA
CTCTGGCTCCTGGCTCTTCAGCCTCATGGCCTCCAGGGCAGAAAAGAGCGGAAATCC
AAGAACTCTTAGACTCTGACCAGCCAGGGCAGGGCAGAGAACGAGCACTTCTAGGCTTGC
TTACTCTACAAGGGACAGTTGCAATTGTTGAGACTTTAATGGAGATTGCTCTACAAGTGGG
AAAGACTGAAGAAAACACATCTGTCAGATCTGCTGGCAGAGGACAATCAAAACGACAACA
AGCTTCTTCCCAGGGTGAGGGAAACACTTAAGGAATAATGGAGCTGGGTTAACACT
AAAAAACTAGAAATAAACATCTCAAACAGTAAAAAAAAAAGGGCGGCCGACTCTAG
AGTCGACCTGCAGAAGCTTGGCCGCACTGGCCAATTGTTATTGAGCTTATAATGGTTAC

FIGURE 56

MDFITSTAILPLLFGLGVFGLFRLLQWVRGKAYLRNAVVVITGATSGLGKECAKVFYAAGA
KLVLCGRNGGALEELIRELTASHATKVQTHKPYLVTFDLTDGAI
VAAAEEILQCFGYVDIL
VNNAGISYRGTIMDTVDVKRVMETNYFGPVALTKALLPSMIKRRQGHIVAISSIQGKMSI
PFRSAYAASKHATQAFFDCLRAEEMEQYEIEVTVISPGYIHTNLSVNAITADGSRYGVMDTTT
AQGRSPVEVAQDVLAAGKKKDVLILADLLPSLAVYLRTLAPGLFFSLMASRARKERKSNS

Signal sequence:

amino acids 1-21

Transmembrane domain:

amino acids 104-120, 278-292

N-glycosylation site.

amino acids 228-232

Glycosaminoglycan attachment site.

amino acids 47-51

Casein kinase II phosphorylation site.

amino acids 135-139, 139-143, 253-257

Tyrosine kinase phosphorylation site.

amino acids 145-153, 146-153

N-myristoylation site.

amino acids 44-50, 105-111, 238-244, 242-248, 291-297

Amidation site.

amino acids 265-269

Prokaryotic membrane lipoprotein lipid attachment site.

amino acids 6-17

FIGURE 57

FIGURE 58

MKFLLDILLPLLLIVCSLESFVKLFIPKRRKSVTGEIVLITGAGHGIGRLTAYEFAKLKSK
LVLWDINKHGLEETAACKGLGAKVHTFVVDCSNREDIYSSAKKVAEIGDVSILVNNAAGVV
YTSDLFATQDPQIEKTFEVNVLAHFWTTKAFLPAMTKNNHGHIVTVASAAGHVSVPFLLAYC
SSKFAAVGFHKTLTDELAALQITGVKTTCLCPNFVNTGFIKNPSTSLGPTLEPEEVVNRLMH
GILTEQKMIFIPSSIAFLTTLERILPERFLAVLKRKISVKFDAVIGYKMKAQ

Signal sequence:

amino acids 1-19

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 30-34, 283-287

Casein kinase II phosphorylation site.

amino acids 52-56, 95-99, 198-202, 267-271

N-myristoylation site.

amino acids 43-49, 72-78, 122-128, 210-216

FIGURE 59

CCACCGCGTCCCGGGACCGCTGGGTCGACTAGTTCTAGATCGCAGCGGCCGCCGCCGCT
AGGGAGGAGCACCGACTGCGCCGACCCCTGAGAGAATGGTGTGCCCCATGTGGAAGGTGATTG
TTTCGCTGTCCTGTTGATGCCTGGCCCTGTGATGGGCTTTGCTCCTATAACAGAAAGT
GTTCCCATGCCACCTAAGGGAGACTCAGGAACAGCATTATTCACCCCTTACATTGAAGC
TGGGAAGATCCAAAAGGAAAGAGAATTGAGTTGGCTGGCCCTTTCCAGGACTGAACATGA
AGAGTTATGCCGCTTCTCACCGTGAATAAGACTTACAACAGCAACCTTCTCTGGTTC
TTCCCAGCTCAGATAACGCCAGAAGATGCCAGTAGTTCTCTGGCTACAGGGTGGCCGG
AGGTTCATCCATTTGGACTCTTGTGGAACATGGGCTTATGTGTCACAAGTAACATGA
CCTGCGTGCAGAGAGACTTCCCTGGACCACACGCTCTCATGCTTACATTGACAATCCA
GTGGGACAGGCTTCAGTTACTGATGATAACCCAGGATATGCACTGATGAGGACATGT
AGCACGGGATTTATACTGCACTAACTCAGTTTCCAGATAATTCTGAATAAAAAATA
ATGACTTTTATGTCAGTGGGAGTCTTATGCAAGGAAATATGTGCCAGCATTGACACCTC
ATCCATTCCCTCAACCCCTGTGAGAGGAGATCACCTGAACCGGATTGCTATTGGAGA
TGGATATTCTGATCCCGAATCAATTATAGGGGCTATGCGAATTCTGTACCAAATTGGCT
TGTGGATGAGAAGAAAAAAAGTACTTCCAGAACGGACTGCCATGAATGCAAGAACACATC
AGGAAGCAGAACACTGGTTGGCCCTTGAAATACTGGATAAAACTACTAGATGGCAGCTTAAC
AAGTGATCTCTTACTTCCAGAATGTTACAGGATGTAGTAAATTACTATAACTTTTGGGT
GCACGGAACCTGAGGATCAGCTTACTATGTGAAATTGGTCACTCCAGAGGTGAGACAA
GCCATCACGTGGGAATCAGACTTTAATGATGGAACCTAGTTGAAAAGTACTTGGAGA
AGATACTACAGTCAAGTCACTGTTAAGGATGAAATCATGAATAATTATAAGGTTCTGA
TCTACAATGGCAACTGGACATCATCGTGGCAGCTGCCCTGACAGAGCCTTGTGGC
ATGACTGGAAAGGATCCCGGAATACAAGAAGGCAGAAAAAAAGTTGGAGATCTTTAA
ATCTGACAGTGAAGTGGCTGGTACATCCGGCAAGCGGGTGACTCCCATCAGGTAAATTATC
GAGGTGGAGGACATATTTCACCTATGACCAAGCCTCTGAGAGCTTTGACATGATTAATCGA
TTCATTATGGAAAAGGATGGATCCTTATGTTGGA**TAA**ACTACCTCCAAAAGAGAACAT
CAGAGGTTTCATTGCTGAAAAGAAAATCTAAAAACAGAAAATGTCATAGGAATAAAAAAA
TTATCTTTCATATCTGCAAGATTTCATCAATAAAATTATCCTGAAACAAGTGAGC
TTTGTGTTGGGGAGATGTTACTACAAAATTAAACATGAGTACATGAGTAAGAATTACA
TTATTTAACTTAAAGGGATGAAAGTATGGATGATGTGACACTGAGACAAAGATGTATAATGA
AATTTTAGGGTCTTGAATAGGAAGTTTAAATTCTTCTTAAGAGTAAGTGAAGGTGCACTTG
TAACAAACAAAGCTGTAACATCTTCTGCAATAACAGAAGTTGGCATGCCGTGAAGGT
GTTGGAAATATTATTGGATAAGAATGCTCAATTATCCAAAATAATGGATGAAGCTATAA
TAGTTTGGGGAAAAGATTCTCAAATGTATAAAGCTTAAAGACAAAAGAATTCTTGAATAA
AAAATATTATATAAAAGTAAAAA

FIGURE 60

MVGAMWKIVSLLMPGCDGLFRSLYRSVSMPPKGDSGQPLFLTPYIEAGKIQKGRELSL
VGPFPGLNMKSYAGFLTVNKTYNSNLFFWFFPAQIQPEDAPVVLWLQGGPGGSSMFGLFVEH
GPYVVTNSNMTLRDRDFPWTTTLSMLYIDNPVGTGFSFTDDTHGYAVNEDDVARDLYSALIQF
FQIFPEYKNNDFYVTGESYAGKYVPAIAHLIHSLNPKREVKINLNGIAIGDGYSDPESIIGG
YAEFLYQIGLLDEKQKKYFQKQCHECIEHIRKQNWFEEAFEILDKLLDGDLTSDPSYFQNVTG
CSNNYNFLRCTEPEDQLYYVKFLSLPENVRAQIAHVGNTFNDGTIVEKYLREDTVQSVKPWL
EIMNNYKVLIYNGQLDIIVAAALTERSLMGMDWKGSQBYKKAEKVVWKIFFKSDSEVAGYIRQ
AGDFHQVIIRGGGHILPYDQPLRAFDMINRFIYGKGWDPYVG

Signal sequence:

amino acids 1-22

N-glycosylation site.

amino acids 81-85, 132-136, 307-311, 346-350

Casein kinase II phosphorylation site.

amino acids 134-138, 160-164, 240-244, 321-325, 334-338, 348-352,
353-357, 424-428

Tyrosine kinase phosphorylation site.

amino acids 423-432

N-myristoylation site.

amino acids 22-28, 110-116, 156-162, 232-238

Serine carboxypeptidases, serine active site.

amino acids 200-208

Crystallins beta and gamma 'Greek key' motif signature.

amino acids 375-391

FIGURE 61

CGAGGGCTTCCGGCTCCGGAAATGGCACATGTGGGAATCCCAGTCCTGTTGGCTAACACAT
TTTCCCTTCTAACAGTCTAACAGCTGTTCAACAGCTAGTGATCAGGGGTTCTCTT
GCTGGAGAAGAAAGGGCTGAGGGCAGAGCAGGGCACTCTACTCAGGGTACCGAGCTCTT
CCTCTGTGATAACAGAGCATGAGAAAGTGAAGAGATGCAGCGGAGTGAGGTATGGAAG
TCTAAAATAGGAAGGAATTTCGAGAATTCAGACTCTGGGAGCAGTTGACCTGGAGAGC
CTGGGGGAGGCCCTGCCAACAAAGCTTCAAACAGGGAGCAGCTTCAACTGGGCTGGGAT
AAGACGTGCGGCTAGGATAGGGAAGACTGGGTTAGTCTTAATATCAAATTGACTGGCTGG
TGAACCTCAACAGGCTTTAACCTCTGTGGAGATGAAACAGATGGCTTAAGGGGCAAGAAA
TAGAGATGCTTGTAAAATAAAATTAAAAAGCAAGTATTATAGCATAAAAGGTAGA
GACCAAAATAGATAACAGGATCTCTGAACATTCTAACAGGGAGAAAGTATGTTAAAATA
GAAAACCAAAATGCAAGAGGAGAGCTCACAGAGCTAAACCAGGATGGGGACCTGGGTC
AGGCCAGCTCTTGCCTCTCCGGAAATTATTTGGTCTGACCAACTCTGCCCTGTGTTT
CGAACATATGTAAGGGGCAACGGGGAGGTGGAGCAGATGAGCACACAGGAGCCGTCT
CCTCACCGCCGCCCTCTCAGCATGGAACAGAGGAGCCCTGGCCCCGGCCCTGGAGGTGG
ACAGCCGCTCTGTTCTGCTCTAGTGGTCTGGTCTGGCCCCCAGCAGCCGGC
ATGCCCTCAGTCAGCACCTTCACCTCTGAGAACATGTCAGTGGACCTTCACCAACTTGACCGT
CCACCAAGGGACGGGGGGCGCTATGTGGGGGCCATCAACGGGTCTATAAGCTGACAGGCA
ACCTGACCATCCAGGTGCTCATAGACAGGGCAGAAGAGGACAACAAGTCTCGTAAACCG
CCCTCATCTGTCAGCAGGGAGCTGGAGCAGGAAAGTCAAAACATGTCAAACAGCTGCT
CATCATTGACTACTCTGAGAACCCCTGTGGGCTGTGGGAGGCTCTACCAAGGGGCTGCA
AGCTGTCGGGCTGGATGACCTCTCATCTGGTGGAGCATCCCACAAGAAGGAGCACTAC
CTGTCAGTGTCAACAAAGACGGGACCATGTCAGGGGTATTGTCGCTCTGAGGGTGAGGA
TGGCAAGGCTTCATCGGCCACGGCTGTGGATGGGAAGCAGGATTAACCTTCCGACCCCTGTCCA
GCCGGAAAGCTGCCCGAGACCTCTGAGTCTCAGGCTCATCGACTATGAGCTACACAGCGAT
TTTGTCTCTCTCATCAAGATCCTTCAGACACCCCTGGCCCTGGTCTCCACTTTGACAT
CTTCATACATCTACGGCTTGCTAGGGGGCTTGTCTACTTCTACTGTCCAGGCCGAGA
CCCTGAGGGTGTGGCCATCAACTCCGCTGGAGACCTCTTACACCTCACGTCATGTGCG
CTCTGCAAGGATGACCCCAAGTCCACTCATCGTCCTGCCCTCGCTGACCCGGC
CGGGGTGGAAATACCGCTCTGAGCTGTCTACCTGGCAAGGCCCTGGGACTCACTGGCC
AGGGCTTCATATCACCGAGGAGCTGACTCTTGCATCTTCTCAAAAGGGCAGAG
CAGTATCACCAACCGCCCGATGACTCTGCCCTGTGCTCTCCCTATCCGGGCACTT
GCAGATCAAGGAGGCCCTGCAACTCTGCTACAGGGCGAGGGCAACCTGGAGCTCAACTGGC
TGCCTGGGAAGGAGCTTCACTGGAGGCTTACACTTCCAGTGGAGGGCTGACCCCTGACACAC
CTGGACATCAACCAGCCCTGGAGGCTTACACTTCCAGTGGAGGGCTGACCCCTGACACAC
CAGCAGGGACCGATGACCTCTGTGGCTCTACGTTAACAGGCTACAGCGTGGTTT
TGGGGACTAAGAGTGTGCAAGGTAAAAAGGTAAAGAGTCTATGAGTTGAGATGCTCAATGCC
ATTCACTCTCTCAGCAAAGACTCTGTGGAGGTTAGCTATTGGTGGAGATTAACTATAG
GCAACTTATTTCTGGGAAACAAGGTGAATGGGGAGGTTAGAAGGGTTAATTTCG
ACTTAGCTTCTAGCTACTCTCCAGCCATCAGTCATTGGTATGTAAGGAATGCAAGCGT
TTCAATATTCACCTTAAGAAAAACTTTAAGAAGGTACATCTGCAAAAGCAAA

FIGURE 62

MGTLGQASLFAPPGNYFWSDHSALCFAESCEGQPGKVEQMSTHRSRLLTAAPLSMEQRQPWP
RALEVDSDRSRVVLLSVVVVLLAPPAAGMPQFSTFHSENRDWTFNHLTVHQGTGAVYVGAINRV
YKLGNLTIQVAHKTGPEEDNKSRYPPLIVQPCSEVLTLNNVNKLIIIDYSENRLLACGSL
YQGVCKLLRLDDLFLILVEPSHKKEHYLSSVNKTGTMGYVIVRSEGEDGKLFIGTAVDGKQDY
FPTLSSRKLPRDPESSAMLDYELHSDFVSSLIKIPSDTLLALVSHFDIFYIYGFAASGGFVYFL
TVQPETPEGVAINSAGDLFYTSRIVRLCKDDPKFHSYVSLPFGCTRAGVEYRLLQAAYLAKP
GDSLAQAFNITSQDDVLFAIFSKGKQYHHPPDDSALCAFPIRAINLQIKERLQSCYQGEQN
LELNWLGLGKDVCQCTKAPVPIDDNFCGLDINQPLGGSTPVEGLTLYTTSRDRMTSVASYVYNG
YSVVFVGTSGKLKKVRYEFRCNSAIHLLSKESSLLEGSYWWRFNYRQLYFLGEQR

Signal sequence:
amino acids 1-32

Transmembrane domain:
amino acids 71-87

N-glycosylation site.
amino acids 130-134, 145-149, 217-221, 381-385

Casein kinase II phosphorylation site.
amino acids 139-143, 229-233, 240-244, 291-295, 324-328, 383-387,
384-388, 471-475, 481-485, 530-534

N-myristoylation site.
amino acids 220-226, 319-325, 353-359, 460-466, 503-509

FIGURE 63

FIGURE 64

MTTWSLRRRPARTLGLLLLVLGFLVLRRLDWSTLVPLRLRHRQLGLQAKGWNFMLEDSTFW
IFGGSIHYFRVPREYWRDRLLKMKACGLNTLTTYVWPWNLHEPERGKFDFSGNLDLEAFVLMA
ABIGLWVLIRPGPYICSEMDLGGLPSWLQDPMRLLRTTYKGFTEAVIDLYFDHLMRSVVPLQ
YKRGGPIIAVQVENEYGSYNKDPAYMPYVKKALEDRGIVELLTSNDKDGLSKGIVQGVLAT
INLQSTHELQLTTFLFNVQGTQPKMVMEYWTGWFDSWGPHNIILDSSEVLKTVSAIVDAGS
SINLYMFHGCTNFGFMNGAMHFHDYKSDVTSYDYDAVLTEAGDYTAKYMKL RDFFGSISGIP
LPPPPDLPKMPYEPLTPVLYSLWDALKYGEPIKSEKPINMENLPVNGGNGQSFGYIYE
TSITSSGILSGHVVHDRGQVFVNTVSIGFLDYKTTKIAVPLIQGYTVLRILVENRGRVNNGEN
IDDKRKGLIGNLYLNDSPPLKNFRIYSLDMKKSFFQRFGLDKWXSLPETPTLPAFFLGSLSIS
STPCDTFLKLEGWEKGVVFINQNLGRYWNIGPQKTLYLPGPWLSSGINQVIVFEETMAGPA
LQPTETPHLGRNQYIK

Signal sequence:

amino acids 1-27

Casein kinase II phosphorylation site.

amino acids 141-118, 253-257, 340-344, 395-399, 540-544, 560-564

N-myristoylation site.

amino acids 146-152, 236-242, 240-246, 244-250, 287-293, 309-315,
320-326, 366-372, 423-429, 425-431, 441-447, 503-509, 580-586

FIGURE 65

GGGGACGCGGAGCTGAGAGGCCGGCTAGCTAGGTGAGGGTGGACGGGTCGGCAGGACC
CTGGTGGAGGTTCTACTTGGCCCTCGGGGGCTAAGACGCAGGACCTACGCAAAGG
GGACAAAGCGGGCTCGGCCGAGGCCCAAGGACCTCACATCCCCTGAGGAGGAATC
CGACACGTGAGCTGCTGCCGCTCAGACTAGAGGAGGCCGTAAACGCCATGGCTCC
AAGAACGCTGCTCTGCCCTCGTCCCTGCTGCCGCTCAGACTAGAGGAGGCCGTAAACGCCATGGCTCC
GGCAGACACTCGCTCGTCTCGTAGTGATAGGGTCATGACCGTTCTCTTAGACGGGCC
CGTCCGCTATGCTGCTGCCAGCCTGCACTACTTCGGTACCGCGGGTCTGGCCGAC
CGCCTTTGAAAGATGCCAGGCCCTAACCGGCTAACAGTTTATGTCGCCCTGGAACTA
CCACGAGCACAGCCTGGGTCTATAATTGAGCAGGGGACCTATTGCCCTTCTGA
ATGAGGCAGCTAGCGAACCTGTTGGTCATACTGAGACGAGGACCTACATCTGTCAGAG
TGGAGATGGGGCTCCCATCTGTTGCTCGAAAACCTGAAATTCTAAGAACCTC
AGATCCAGACTTCTTGCAGTGACTCTGGTCAAGGCTTGTGCTGCCAACATATATC
CATGGCTTATCACAAATGGGGCACACATTAGCATTCAAGGTGAGAATGAAATATGGTAGC
TACAGAGCTGTGACTCTAGCTACATGAGGCACTTGGCTGGCTTCCGTGCACTGCTAGG
AGAAAGATCTGCTCTCACACAGATGGGCCAGTCAAGGACTCAAGTGTGGCTCCCTCCGG
GACTCTATACCATCTAGATTTGGCCAGTGCACAACTGACCAAAATCTTACCTGCTT
CGGAAGTGAACCCATGGGCCATTGGTAACACTCTGAGTACTACACAGGCTGGCTGGATT
CTGGGGCCAGAACATCCTCACACGGTCTGGTCAAGGCTGTAACCAAGGACTAGAAACATGC
TCAAGTGGGGCCAGTGTGACATGGTACATGTCATGGGAGGTACCAACTTGGATATTGG
AATGGTGCAGATAAGAAGGGACCTTCTCCGATTACTACAGCTATGACTATGATGCACC
TATATCTAAGCAGGGACCCACACCTAACGCTTTTGCTCTCGAGATGTCACTAGCAAGT
TCCAGGAAGTTCTTGGGACCTTACCTCCCCGAGCCCAAGAGATGCTTGGACCTG
ACTCTGCACCTGGTGGGCAATTACTGGCTTCTGAGACTGCTTGCCTCCGGTGGCCCAT
TCATTCAATCTGCCAATGACCTTGGAGGCTGCAAGCAGGACCATGGCTCATGGTAC
GAACCTATATGCCATACCATTTTGAGGCAACACCATCTGGGTGCCAAATAATGGAGTC
CATGACCGTCTCTATGATGGTGGATGGGTCTCAGGGTGTGAGGCGAAATATGAG
AGACAAACTATTGGACGGGAAACTGGGCTCAAATGGGATATCTGGTGGAGAACATGG
GGAGGCTAGCTTGGGCTAACAGCAGTGACTCAAGGGCTGTTGAAGCCACCAATTCTG
GGCAAAACATCTTACCTGGGCAATTCTGGGCTAACATTGGGATTAACCTTGTGAAGT
GTGGTTCCCTCCAGTTGCCAAATGGGCAATTCTGGGCTAACATTGGGCAACATTCT
ACTCCAAAACATTCCAATTGGCTAGTGGGACACATTCTATATCTACCTGGATGG
ACCAAGGGCCAAGTCTGGATCAATGGGTTAACCTGGGCGTACTGGACAACAGGGCC
ACACAGACCTACCTGCCAACATTCTGGTGGGACCTAACAAAATTAA
CATTGCTGAAACTAGAAGATGTACCTCTCACAGGCAACATATCAATTCCCTTCTGAG
CTCAATAGCAACTACTTGCACAGGACACATATCAATTCCCTTCTGAGCTGATAACACTGAG
TGCTCTGAACCAATGGGAGTTAAGTGGGCACTGAAAGGTAGGCCGGCATGGTGCTCATGC
CTGTAATCCAGCACTTGGGAGGCTGAGACGGGTGGATTACCTGAGGTCAAGGACTTCAAGA
CCAGCCTGCCAACATGGTGAACACCGTCTCAACTAAAAATACAAAATTAGCCGGCGTG
ATGGTGGGCACCTCAATCCAGCTACTTGGGAGGCTGAGGGCAGGAGAATTGCTTGAATCC
AGGAGGAGAGGTTGCACTGAGTGGAGGTTGTAACACTGCACCTCCAGCCTGGCTGACAGTGA
GACACTCCATCTAAAAAAAAAA

FIGURE 66

MAPKKLSCLRSLLLPLSLTLLPQADTRSFVVDRGHDRFLLDGAPFRYVSGSLHYFRVPRVL
WADRLLKMRWSGLNAIQFYVPWNYHEPQPGVYNFNGSRDLIAFLNEAALANLLVILRPGPYI
CAEWEMGGGLPSWLLRKPEIHLRTSDPDFLAADVDSWFKVLLPKIYPWLYHNGGNIISIQVE
YGSYRACDFSYMRHLAGLFRALLGEKILLFTTDGPEGLKCGSLRGLYTTVDFGPADNMTKIF
TLLRKYEPHGPLVNSEYYTGWLWDYWGQNHSRSVSATKGLENMLKLGASVNMYMFHGGTNF
GYWNGADKKGRFLPITTSDYDAPISEAGDPTPKLFALRDVISKFQEVPGLPLPPSPKMM
GPVTLHLVGHLAFLDLLCPRGPIHSILPMTFEAVKQDHGFLYRTYMTHTIFEPTFWVPN
NGVHDRAYVMVGVFQGVVERNMRDKLFLTGKLGSKLDILVENMGRSLFGSNSSDFKGLLKP
PILGQTILTQWMMFPLKIDNLVKWWFPLQLPKWPYQPQAPSGPTFYSKTFPILGSVGDTFLYL
PGWTKGQVWINGFNLGRYWTQKQGPQQTLYVPRFLFPFRGALNKITLLELEDVPLQPQVQFLD
KPILNSTSTLHRTHINSLSADTLSASEPMELSGH

Signal sequence:

amino acids 1-27

N-glycosylation site.

amino acids 97-101, 243-247, 276-280, 486-490, 625-629

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 4-8

Casein kinase II phosphorylation site.

amino acids 148-152, 234-238, 327-331, 423-427, 469-473, 550-554,
603-607, 644-648

Tyrosine kinase phosphorylation site.

amino acids 191-198

N-myristoylation site.

amino acids 131-137, 176-182, 188-194, 203-209, 223-229, 227-233,
231-237, 274-280, 296-300, 307-313, 447-453, 484-490

FIGURE 67

GCTTGAAACACGCTGCAAGCCAAAGTTGAGCATCTGATTGGTTATGAGGTATTCAGTGC
ACCCACAATATGCCTTACATGTTGAAAAGCTCTCATCAGTTACATATCATTATTTGTGT
TTATGGCTTATCTGCCTCTACACTCTCTGGTTATTCAAGGATACCTTGAAGGAATATT
CTTCGAAAAAGTCAGAGAAGAGCAGTTAGTGCACATCCAGATGTCAAAACGATTT
GCGTTCTCTTCACATGGTAGACCGAGTATGACCAGCTATATTCCAAGCCTTGGTGTGTT
CTTGTCAAGAGTTAGTCAAATAACTTAGGGAAATTAGTGTGAACCAGTGCATCTTCATG
AAAAACTCAGGCAGCACATTCAACGCCAGGACAAGCAGGAGTTGCATCTTCATG
CTGTCGGGGTGGCCGATGCTGCTTGACCTCACAGACCTGGATGTGCTAAAGCTTGAAC
AATTCCAGAAGCTAAAATTCTGCTAACAGATTCTCAATGACTAACCTCAAGAGCTCACC
TCTGCCACTGCCCTGCAGAAAAGTTGAACAGACTGCTTTAGCTTCTCGCGATCACTTGAGA
TGCCCTCAGTGAAAGTTCACTGATGTTGCTGAAATTCTGCTGGGTGATTTGCTAAAAAA
CCTTCGAGAGTTGTACTTAATAGCAATTGAACTCTGAAAACAATAAGATGATAGGACTTG
AAATCTCTCGAGAGTTGCGGCACCTTAAGATTCTCACGTGAAGGCAATTGACCAAAGTT
CCCTCCAACATTACAGATGTGGCTCCACATCTAACAAAGTTAGTCATTCAATGACGGCAC
TAAACTCTGGTACTGAACAGCCTAACAGAAAATGATGAATGTCGCTGAGCTGGAACCTCAGA
ACTGTGAGCTAGAGAGAACATCCACATGCTATTTCAGCCTCTCAATTACAGGAACTGGAT
TTAAAGTCAATAAACATTGCAACATTGAGGAAATCATCAGTTCCAGCATTAAAAGCACT
GACTTGTAAAATTATGGCATAACAAAATTGTTACTATTCTCCCTCTATTACCCATGTCA
AAAACCTGGAGTCACTTATTCCTAACACAAGCTGAATCCTTACAGTGGCAGTATT
AGTTTACGAAAACTCAGATGCTTACATGAGCTACAACAAACATTCAATGATTCAATAGA
AATAGGATTGCTCAGAACCTGCAGCATTGCAATATCACTGGAACAAAGTGGACATTCTGC
CAAAACAATTGTTAACATGCAAAAGTTGAGGACTTGAATCTGGGACAGAACTGCATCACC
TCACCTCCAGAGAAAAGTTGGTCAGCTCTCCAGCTCACTCAGCTGGAGCTGAAGGGAACTG
CTTGGACCGCCTGCCAGCCCAGCTGGGCCAGTGTGGATGCTCAAGAAAAGCAGGGCTTGTG
TGGAAAGATCACCTTTTGATACCCGCCACTCGAAGTCAAGAGGCAATTGAATCAAGACATA
AATATTCCCTTGCAATGGGATTAAACTAAGATAATATATGCACAGTGATGTGCAGGAAC
AACTTCTAGATTGCAAGTGCTCACGTACAAGTTATTACAAGATAATGCATTAGGAGTAG
ATACATCTTTAAAATAACAGAGAGGATGCATAGAAGGCTGATAGAAGACATAACTGAAT
GTTCAATGTTGAGGTTAAAGTCATTCAATTCAACATTGTTAAATTGTTGTAACCTGGAT
AAAGGGAGGAAAATTATAACTAACTATCTGGTTCTTTAAATTGTTGTAACCTGGAT
GCTGCCGCTACTGAATGTTACAAATTGCTGCCACTGCTAAAGTAAATGATTAATTGACATT
TTCTTACTAAAAAAAAAAAAAA

FIGURE 68

MAYMLKKLLISYISIICVYGFICLYTLFWLFRIPLKEYSFEKVREESSFSDIPDVKNDFAFL
LHMVDQYDQLYSKRGFGVFLSEVSENKLREISLNHEWTFEKLRQHISRNAQDKQELHLFMLSG
VPDAVFDFLDLVKLELIPEAKIPAKISQMTNLQELHLCHCPAKVEQTAFSFLRDHLRCLH
VKFTDVAEIPAWVYLLKNLRELYLIGNLNSENNKMIGLESRLERLRLKILHVKSNLTKVPSN
ITDVAPHLTKLVIHNDGTKLVLNSLKKMMNVAEELQNCELERIPIHAIFSLSNLQELDLKS
NNIRTIEEEISFQHLKRLTCLKLWHNKIVTIPPSITHVKNLESLYFSNNKLESLPVAVFSLQ
KLRCLDVSYNNISMIPIEIGLLQNLQHLHITGNVKVDILPKQLFKCIKLRTLNLGQNCITSPL
EKVGQLSQLTQLELKGNCLDRLPAQLGQCRMMLKKSGLVVEDHLDFTLPLEVKEALNQDINIP
FANGI

Signal sequence:

amino acids 1-20

N-glycosylation site.

amino acids 241-245, 248-252, 383-387

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 326-330

Casein kinase II phosphorylation site.

amino acids 48-52, 133-137, 226-230, 315-319, 432-436, 444-448

Tyrosine kinase phosphorylation site.

amino acids 349-355, 375-381

N-myristoylation site.

amino acids 78-84, 124-130, 212-218, 392-398

FIGURE 69

CCCCACCGGTCCGGCTTCTCTGGACTTTCACTTCCATTGACAAACTGACTTTTTATTTCT
TTTTTTCATCTCTGGGCAGCTTGGAATCCTAGGCCCTGGAAAGACATTGTGTTTACACATAAGGAT
CTGTTTGGGGTTCTCTCTCCCTGACATTGCAATTGCTTAGGTCTAGGTTGTTGGGGAGGGACACCTG
GTCAGTCTGCTGCACTTATCTGCCCTAGGTACATCGAAGTCTTTCACCTACAGTGAATTGCTG
ATCGCTGTGATCTCTGGCCCTGCTCTGCTGATAGTTGCTGCTGCTGCTTACTCAAAATACAAAC
GCGCTAAAAGCTGCAAGGAACCTGAGACTGTGCTGCTGCTGCTGCTGCTGCTGCTGCTG
AACAGCCAGGCAAACCATTECCACCGAGCTTGTCTGCCCTGAGTGTGAAGGATATAGAATGTG
AGTTTGTATTCCTGCCACCTTGTGTTGGACATAATGGGGCCTCTGAGTTAGGAAAGGCTCCCTCTCAA
GCAAGAGCCCTGAAGACTTCAATGATGTCATGGGCCACCTGTGATGTGAGGACAGAGGCTTC
CTCCCCATCAGTTCACTGGAAATAACTCAGTGCCTGCTGGACCAGCTGCTGGAGATCCTACAGAGGCTTC
CACTGCTGCCACCCCTCAGGAAGAGTGTGGGGAGAGAAACCTCAGTGTGGGAATGCTGATAAACCCAGTCA
CACAGCTCTATTCTCACAAATCTACCCCTTGCGCTGCGTGAACGATCTTCTCTGGAGGTGTCAGAAA
GCTGATGTAACACAGAGCTTAAAGCTGAGCTGGCTTAAAGGCTGCCAGCCTTGCCTAAGGAGCTTGT
AGAAGGCTCATGCCATTGACCTCTTCTCTGTGTTGGGGGAGTCACATGGGGAGGTGAGGCAAT
GCAAGCTGCAAGCTAGCTAGGGGGTGCCTAATGGCAGAGACCACAAAGCCATGATCTGCAACTCAATCC
AGTGAAGAAGTGCACCTGGACAATAGAAGCAGAGAAACCCAGTCAAGAATTATCTTCTATGTCAGCTT
GATCCAGATGGAAGGCTGAAAGTGAACAAATTAAGCTTGTGAACTCATCAGTCAATTGAGCTTCAAATAGT
CAAGTCTGCACTTAAACAGCTATGCTTCTCTACTCTCTCTCTCTCTCTCTCTCTCTCT
GGCGTTACCTGGATACCTTGGAGGATCTTCAACAGGCCAATTACCCAAGCCGATCTGAGCTGGCTTAT
TGTGTGTGGCACATCTGGAGGAAAGTAAACGATAAAACTAACTTCAAGGATTTCTGGAAATAGAC
AAACAGTCCAATTGTATTTCTGCCATCTGATGCCCCCTCCACCAACTCTGGCTGATGGACAGCTGT
GGCGTGTGACTCCACCTTGTCAATCAAACTCTGACTCTGACTGTGTTGTTCTACAGATTATGCAAACTT
TACCGGGGATTCTCTGCTTCTACACCTCAATTGTGCAAAACATCACACTCATCTTAACTGTCTCT
GACAGGATGAGGTTTAAATGCAAACTTACCTAGAGGCTTTAACCTTAATGGAAATACACTGCAACTAAA
GACCCAACTTGCAGACAAJAAATTATGAGGTTGAAATTCTGCTTCAACTCTGATCTCAACTCTGA
AAGGTTAGAAGATCAGTCATTACTACACCAATTATACTCCTTCTGATCTCAACTCTGAAGTGTACATCAGA
CGTCAGAAACAACCTCCAGATTATTGTGAAGTGTGAATGGGACATAATTCTACAGTGGAGATAATATACTAA
GAAGATGATGTAAATCAAAAGTCAACTGGGAAATATAACACCCAGCATGGCTTTTGAAATCCAATTCA
TTTGAAGAAGACTATACCTGAAATCACCATTATGTGGATTGTGAACCAAACTCTTGTCAAGTTAGTC
ACCTCAGATCCAATTGTGGTTGTTCTGATACCTGTAGAGGCCCTCCACCTCTGACTTTGCACTCTCAACC
TACAGCCTTAATCAAGAGTGGATGTAGTCAGATGAAACTCTGAAGGTGATCTCCCTATTGGACACTATGGAGA
TTCAGTTAATGCCCTTAAATTCTTGAGAACTATGAGCTCTGTGATCTGCACTGTAAGTTTGATATGT
AGCAGTGCACCACTCTGCCCTGCATCAAGGTGTCTCCAGGACAAACAGACATTCTTCTATATAATGG
AAACAGATTCCATCATAGGGCCCATTCGCTCTGAAAGGGATCGAACAGTGGCAATTAGGATTCAGCAT
GAAACACATGCCGAAGAAACCTCCAAACAGCCCTTCAACAGTGTGCACTCTTCTCATGGTTCTAGCTG
AATGTGGTGACTGTGCGACAATCACAGTGAGGCATTGTAATCACCGGAGCAGACTACAAATACAGAGCTG
CAGAACTTAACTAACAGGTCCAAACCTTAAGTGAGACATGTTCTCCAGGATGCCAAAGGAAATGCTACCTCGT
GGCTACATCATATTGTAATGAGGAAGGCCCTGAAGACTGACACACAGGCCCTGATGTAACAAAAAA

FIGURE 70

MELVRRMLPLTLILSCLAEILTMAEAEGNASCTVSLGGANMAETHKAMIQLQNPSENCTWTI
ERPENKSIIRIIFSYVQLDPDGSCSESENIVFDCGTSSNGPILLGQVCSKNDYVPVFESSSTLT
FQIVTDSARIQRTVFVFFSPNISIPNCGGYLDTLEGSFTSPNYPKPHPELAYCWHIQV
EKDYKIKLNFKEIFLEIDKQCKFDLAIYDGSTNSTGLIGOCVGRVTPTFESSSNSLTVVLS
TDYANSYRGFSASYTSIYAENINTTSLTCSSDRMRVIISKSYLEAFNSNGNNLQLKDPTCRP
KLSNVVEFSVPLNGCGTIRKVEDQSITYTNIIITFSASSTSEVITRQKQLQIIVKCEMGHNST
VEIYYITEDDVIQSQNALGKYNTSMALFESNSFEKTILESPYYVDLNQTLFVQVSLHTSDPN
LVVFLDTCRASPTSDFASPTYDLIKSGCSRDETCKVYPLFCHYGRQFQNAFKFLRSMSSVYL
QCKVLICDSSDHQSRCNQGCVSRSKRDISSYWKTDSDIIGPIRLKRDRSASGNSGFQHETHA
EETPNQPFMSVHLFSFMVLALNVTTVATITVRHFVNQRADYKYQKLQNY

Signal sequence:

amino acids 1-24

Transmembrane domain:

amino acids 571-586

N-glycosylation site.

amino acids 29-33, 57-61, 67-71, 148-152, 271-275, 370-374,
394-398, 419-423

Casein kinase II phosphorylation site.

amino acids 22-26, 108-112, 289-293, 348-352, 371-375, 379-383,
408-412, 463-467, 520-524, 556-560

Tyrosine kinase phosphorylation site.

amino acids 172-180, 407-415, 407-416, 519-528

N-myristoylation site.

amino acids 28-34, 38-44, 83-89, 95-101, 104-110, 226-232

Prokaryotic membrane lipoprotein lipid attachment site.

amino acids 7-18

FIGURE 71

GACGGAAAGAACAGCGCTCCCGAGGCCGCGGGAGGCCTGCAGAGAGGACAGCCGGCCTGCGCCG
GGACAT**TGCGCCCCAGAGCTCCCAGGCTCGCGTCCCGTGTGCTGTTGCTGTC**
TGCTGCCGCCGCCGCCGCTGCCAGGCCACAGCGCACCGCGCTTCGACCCACACTGGGAGTCC
CTGGACGCCGCCAGCTGCCCGTGGTTGACCAAGGCAAGTCCGGCATCTTCATCCACTG
GGGAGTGTTCCTGGCCAGCTCGTAGGAGTGGTTCTGGTGGTATTGGCAAAGGAAA
AGATACCGAAGTATGTGGAATTATGAAAGATAATTACCCCTCTAGTTCAAATATGAAGAT
TTTGGACCACTATTTACGAAAATTTTAATGCCAACACTGGGAGATATTTCAGGC
CTCTGGTGCCAATACATTGCTTAACCTCCAAACATCATGAAGGCTTACCTTGTTGGGGT
CAGAATATTCTGTGGAATGCCATAGTGAGGGGCGCAAGGGGACATTGTCAAGGAA
CTTGAGGTACCCATTAGGAACAGAAGTGCACCTGGCTTTGGACTGTACTATTCCCTTTGTA
ATGGTTTCATCCGCTCTCCTGAGGATGAATCCAGTTCACTCCATAAGCGGCAATTCCAG
TTCTAAGACATTCGCGAGCTCTAGTGTAGTGTAGAACACTATCAGCTGAGGTTCTGTGG
TCGGATGGTGAGGGAGGACCCGGATCAATCTGGAACAGCACAGGCTTCTGGCTGGTT
ATATAATGAAAGCCCAGTCTGGGGCACTAGTAGTCAACATGATCTGGAGCTGTAGCA
TCCTGAAAGCTGGTGTCTTCTATACTCTGAGTGTGTATAACCCAGGACATTTGCCA
CATAAATGGGAAAAGTGCATGACAATAGACAACACTGCTCTGGGCTATAAGGAGGGAAAGCTGG
AACTCTGACTATCTTACAATTGAAAGATTGGTAGAACACTTGTAGAGACAGTTCTGTG
GAGGAAATCTTTGATGAATTGGGCCCACTAGATGGCACCATTCTGTAGTTTGAG
GACCGACTGAGGCAACTGGGCTCTGGCTTAAAGTCAATGGAGAAAGCTATTGAAACCTA
TACCTGGCGATCCAGAATGACACTGTCACCCAGATGTGTGTACACATCCAAGCCTAAAG
AAAAATTAGTCTATGCCATTCTTCTTAAATTGGGCCACATCAGGACAGCTTCTGGCCAT
CCCAAAGCTATTCTGGGGCAACAGAGCTGAAACTACTGGGCTATGGACAGCCACTTAAC
GATTCTTGGGAGCAAATGGCATTATGGTAGAACACTGCCAACAGCTAACCATTCAGATGC
CGTGTAAATGGGCTCTGGCTCTAGCCCTAACTATGTGAT**CTAAAGTGCAGCAGTG**
ATGCTGCAAGTTATGTCTAAGGCTAGGAACATCAGGTGTCTATAATTGTAGCAGATGGAGA
AACCAATGTAAACTGGATAAGAAAATTGGCAGTTCTGGCAGGCTTCCCTTCCACTA
ATTTTCTTAAATTACCCAGTAACTTAACTCTCAGTGCACTTGCCATTAAAGTC
TCTTCACATTGATTGTTCCATGTGTGACTCAAGGGTAGAATTTCACATTAGTAG
CAAGGAATTGCTGGTATTATGGACCCAACTGAAATTTTATGTGTAGGACATATCCCCATG
ATTATATGTTATGCTACACTTAATATGGGATATTCTGGGAAATGCAATTGCTAGTCAAT
TTTTTTCTGCCAACATCATAGTGTATTACAAATCTAGATGGCATAGCCACTACA
CACCTAATGTGTATGGTATAGACTGTGTCTCTAGGCTACAGACATATACAGCATGTTACTG
AATACTGTAGGCAATAGTAAACAGTGATTGTATATCGAAACATATGGAAACATAGAGAAG
GTACAGTAAAATACTGTAAAATGGTCACCTGTATAGGGCACTTACACAGAAATGGAG
CTTACAGGACTGGAAGTGCTGGTGAGTCAGTGAAGTGAAGGCTAGGACATTA
TTGAAACACTGCCAGACGTTATAAAACTGTATGCTTAGGCTACACTACATTATAAAAAAA
GTTTTCTTCTCAATTATAAACTGTAAAGTGTACTGTAACTTACAAACGTTTAATT
TTTAAACCTTTGGCTCTTTGTAATAACACTTGTAAACATAAAACTCATTTGCAAA
ATGTAA

FIGURE 72

MRPQELPRLAFPLLLLLLPPPCPAHSATRFDPTWESLDARQLPAWFDQAKFGIFIHWG
VFSVPSFGSEWFWWYWKKEKIPKYVEFMKDNYPPSFKYEDFGPLFTAKFFNANQWADIFQAS
GAKYIVLTSKHHEGFTLWGSEYSWNWNAIDEKGPKRDIVKELEVPAIRNRTDLRGFLYYSLFEW
FHPLFLEDESSSFHKRQFPVSKTLPLEYELVNYYQPEVLWSDGDGGAPDQYWNSTGFLAWLY
NESPVRGTVVTNDRWGAGSICKHGGFYTCSDRYNPGHLLPHKWENCMTIIDKLSWGYRREAGI
SDYLTIEELVKQLVETVSCGGNLLMNINGPTLDGTISVVFEERLRQVGWSLKVNGEAIYETY
WRSQNNDTVPDVWYTSKPKEKLVYAIFLKWPTSGQLFLGHPKAILGATEVKLLGHGQPLNWI
SLEQNGIMVELPQLTIHQMPCKWGWLALTNVI

Signal sequence:

amino acids 1-28

N-glycosylation site.

amino acids 171-175, 239-243, 377-381

Casein kinase II phosphorylation site.

amino acids 32-36, 182-186, 209-213, 227-231, 276-280, 315-319,
375-375

Tyrosine kinase phosphorylation site.

amino acids 361-369, 389-397

N-myristoylation site.

amino acids 143-149, 178-184, 255-261, 272-278, 428-434

Leucine zipper pattern.

amino acids 410-432

Alpha-L-fucosidase putative active site.

amino acids 283-295

FIGURE 73

AGCAGGGAAATCGGATGTCTGGTTATGAAGTGGAGCAGTGAGTGTGAGCCTCAACATAGT
TCCAGAACCTCCATCGGACTAGTTATTGAGCATCTGCCTCTCATATCACCACTGGCCATC
TGAGGTGTTCCCTGGCTCTGAAGGGTAGGCACGATGGCAGGTGCTTCAGCCTGGTGTG
CTTCTCACTCCATCTGGACCACAGGGCTCTGGTCAAGGCTCTTGCGTGAGAAGAGCT
TTCATCCAGGTGCTAGCAGAAATTATGGGGATCACCCCTTGTGAGCAAAAAGGGCAACAGC
AGCTGAATTTCACAGAACGCTAAGGAGGCCCTGAGGCTGGGACTAAGTTGGCCGGCAAG
GACCAAGTTGAAACAGCCTTGAAGAGCTAGCTTGAACCTTGAGCTATGGCTGGGTTGGAGA
TGGATTCTGGTCACTCTAGGATTAGCCTAACCCAAAGTGTGGGAAAAAATGGGTGGGTG
TCTGATTGGAAAGGTTCCAGTGAGGCCAGCTTGCAGCTATTGTTACAACCTCATCTGAT
ACTTGGACTAACTCTGTCATCCAGAAATTATCACCAACAAAGATCCCATATTCAACACTCA
AACTGCAACAAACACAGAATTATTGTCACTGACAGTACACTCCTCCGTCAGCTTCACTTCA
ACTCTACAAATCTGGCCCTACTACTCTCCAGTGGCTTCACTTCTATTCCACGG
AGAAAAAAATTGATTGTGTCAGAAGTTTATGAAACTAGCACCATGTCAGAAC
TGAACCATTGTTGAAAATAAAGCAGCATTCAAGAATGAAGCTGCTGGGTTGGAGGTGTC
CCACGGCTCTGCTAGTCTGGCTCTCTCTTGTGCTGAGCTGGTCTGGATTTC
TATGCAAAAGGTATGTAAGGCCCTTCTTACAAACAGAATCAGCAGAAC
CGAACAAAAGTAGTAAAGGGAGAAGGCCATGATGCAACCCCTAATGAGGAATCAAAGA
AAACTGATAAAAACCCAGAACAGAGTCCAAGAGTCCAAGAACAAACTACCGTGCATGCTGGAA
GCTGAAGTTAGATGAGAACGAGAACGAGACACCTGAGGCTGGTTCTTCATGCTCC
TTACCCCTGCCCAAGCTGGGAAATCAAAGGCCAAAGAACAAAGAACAGGAAAGTCACCC
GGTCTCTAACTGGAATCAGCTCAGGACTGCCATTGGACTATGGAGTGCACCAAGAGAATGC
CCTCTCTTATGTAACCTCTGATCCTATCCTCCACCTCCAAAGCTTCCCAGGCC
TTTCTAGCCTGGTATGCTCTAAATATCCACTGGGAGAACGGAGTTTGCAAGTGC
GGACCTAAACATCTCATCAGTATCCAGTGGTAAAAGGCCCTCTGGCTGCTGAGCTAGG
TGGGTTGAAAGCCAAGGAGTCAGTGGACCAAGGCTTCTACTGATTTCGAGCTCAGAC
CCTCTCAGCTCTGAAGAACACGTATCCACCTGACATGCTCTTGAGCCCCGTA
AGAGCAAAGAACGAGAAAAGTTAGCCTGAAAGGCCATTCTATAACTTGTG
ACCTAATCTGTAAGCTAAATAAAGAACATGACAAAGGCTGAGGATACGACAGTACACT
GTAGCAGGGACTGTAACACAGAACAGGCTAAAGTGTCTCTGAAACACATTGAGITGGA
ATCACTGTTAGAACACACACACTTACTTTCTGGTCTCTACACTGCTGATATTCTCT
AGGAAATATACCTTCTAACAGTAACAAAATAAAACTCTTAAATTCTATTCTATCTGA
GTTACAGAACATGATTACTAACGGAGATTACTCAAGTAATTGTTAAAAGTAATAAAATTCA
ACAACACATTGCTGAATGCTACTATATGTCAGTGCAAGGTATACACTCTGTAAT
TGAATTATTCTCTAACAAAATTGACATAGTGAAGCTATCTGGGAAGCTATTCTCA
GTTTGATATTCTAGCTTATCTACTTCCAAACTAATTCTATTCTGAGACTAATCTT
ATTCACTTCTCTAACATGGCAACCCATTATAACCTTAAATTATTAAACATACCTAACAG
TACATTGTTACCTCTATACAAAGCACATTAAAAGTGCCTTAAACAAATGTACTCA
GCCCTCTTCTCAACAGAAGGGACTGAGAGATGCAGAAATATTGTGACAAAAATTAA
AGCATTAGAAAATT

FIGURE 74

MARCFSLVLLLTSIWTRRLVQ GSLRAEELSIQVSCRIMGITLVSKKANQLNFTEAKEACR
LLGLSLAGKDQVETALKASFETCSYGWVGDFVVISRISP NPKCGKNGVGLIWKVPVSRQF
AA YCYNSSDTWTNSCIPEIITKDP IFNTQTATQTTEFIVSDSTYVASPYSTIPAPTTPP
APASTSIPRRKKLICVTEVFMETSTMSTETEPFVENKA AFKNEAAGFGGVPTALLVLALLFF
GAAAGLGFCYVKRYVKAFFTNKNQQKEMIETKVVKEEKANDSNPNEESKTDKNPEESKSP
SKTTVRCLAEV

Signal sequence:

amino acids 1-16

Transmembrane domain:

amino acids 235-254

N-glycosylation site.

amino acids 53-57, 130-134, 289-293

Casein kinase II phosphorylation site.

amino acids 145-149, 214-218

Tyrosine kinase phosphorylation site.

amino acids 79-88

N-myristoylation site.

amino acids 23-29, 65-71, 234-240, 235-239, 249-255, 253-259

FIGURE 75

AGATGGCGGTCTGGCACCTCTAATTGCTCTCGTGTATTCGGTGCACGACTTTACGATGG
CTCGCCCAACCTTACTACCTTCTGTCGGCCCTGCTCTGCTGCCCTACTCGTGAGGAA
ACTGCGCCGCTCTGCCACGGCTGCCACCCAACGCGAAGACGTAACCGTGTGACTTTG
ACTGGAGAGAAGTGGAGATCCTGATGTTCTCAGTGCCTGTGATGATGAAGAACCGCAGA
TCCATCACTGTGGAGCAACATATAGGCAACATTTCATGTTAGTAAAGTGGCCAACACAAT
TCTTTCTTCCGCTTGGATATTGCATGGGCCACTTACATCACACTGCTAGTGTCC
TGATGACGTGCAAACCCCCCTATATATGGGCCCTGAGTATATCAAGTACTCAATGATAAA
ACCATTGATGAGGAACCTAGAACGGGACAAGAGGGTCACTGGATTGTGGAGTTCTTGCAA
TTGGCTAACTGCAATCATTGCCCCATCTATGCTGACCTCTCCCTAAATACAAC
GTACAGGGCTAAATTGGAGGTGGATGTTGGACGCTATACTGATGTTAGTACGGTAC
AAAGTGAGCACATCACCCCTACCAAGCAACTCCCTACCTGATCCTGTTCCAAGGTGGAA
GGAGGCAATCGGGGGCCACAGATTGACAAGAAAGGACGGCTGTCTCATGGACCTCTCG
AGGAGAATGTGATCCGAGAATTAACTTAAAGCTATACCAGGGCCAAGAAACTATCA
AAGGCTGGAGACAATATCCCTGAGGAGCAGCTGTGGCTCAACCCCCACAGTGTAGA
TGGGAAAACAAGAAGGATAAATAAGATCCTCACTTGGCAGTGCTCCCTGTCATT
CCAGGCTTCCATAACCACAAGCCTGAGGCTGCAGCCTTNATTNATGTTTCCCTTGG
CTGNGACTGGNTGGGGCAGCATGCAGCTCTGATTAAAGAGGCATCTAGGGAAATTGTCAG
GCACCCCTACAGGAAGGGCTGCCATGCTGTGGCCACTGTTCACTGGAGCAAGAAAGAGATC
TCATAGGACGGGGGGAAATGGTTCCCTCAAGCTGGGTAGTGTGTTAAGCTGCTTATC
AGCTATTACAGACATCTCCATGGTTCTCCATGAAACTCTGTGGTTCATCATTCCCTGTTAG
TTGACCTGCACAGCTGGTTAGACCTAGATTTAACCTAAGGTAAGATGCTGGGTATAGAA
CGCTAAGAATTTCCTTCAAGGACTCTGCTTCTTAAGGCCCTCTGGCTTCTGGTC
TTCAATTAAAGTATAAGCTAACCTTGTGCTAGTCTTAAGGAGAAACCTTAAACCACAAAG
TTTTTATCATTGAAAGACAATATTGAAACAACCCCTATTGTGGGGATTGAGAAGGGTGAA
TAGAGGCTTGGAGACTTCCCTTGTGTGGTAGGACTTGGAGGAGAAATCCCTGGACTTAC
TAACCCCTGACATACTCCCCACACCCAGTTGATGGCTTCCGTAAATAAAAGATTGGGATT
TCCTTTG

FIGURE 76

MAVLAPLIALVYSPVRLSRWLAQPYYLLSALLSAFLVRKLPPCHGLPTQREDGNPCDFD
WREVEILMFLSAIVMMKNRRSITVEQHIGNIFMFMSKVANTILFFRLDIIRMGLLYITLCIVFL
MTCKPPLYMGPEYIKYFNDKTIDEELERDKRVTVIVEFFANWSNDCQSFAPIYADLSLKYNC
TGLNFKGKVDVGRYTDVSTRYKVSTSPLTKQLPTLILFQGGKEAMRRPQIDKKGRAVSWTFSE
ENVIREFNLNELYQRACKLSKAGDNIPEEQPVASTPTTVSDGENKKDK

Signal sequence:

amino acids 1-48

Transmembrane domain:

amino acids 111-125

N-glycosylation site.

amino acids 165-169, 185-189

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 154-158, 265-269

Casein kinase II phosphorylation site.

amino acids 51-55, 145-149, 245-249, 286-290, 288-292

N-myristoylation site.

amino acids 188-194, 225-231

Myb DNA-binding domain repeat signature 1.

amino acids 244-253

FIGURE 77

GGACAGCTCGGGCCCCGAGAGCTCTAGCCGTGAGGAGCTGCCTGGGACGTTGCCCTG
GGGCCCAGCCTGGCCGGTCAACCTGGCATGAGGAGATGGCTGTTCTCGTCCCTGGTCCA
TTGCTCTGCTGCCGGCTCTACGGACTGCCCTCTACAACGGCTTCACTACTCCAAACAG
CGCCAACGACCAGAACCTAGGCAACGGTCATGGCAAAGACCTCTTAATGGAGTGAAGCTGG
TGGTGGAGACACCCGAGGGAGACCCCTGTTACCCCTACCAAGGGGCCAGTGTGATCCTGCCCTGC
CGCTACCGCTACGAGGCCGGCTGGTCTCCCGCGCGTGTGCGTGTCAAATGGTGGAAAGCT
GTGGGAGAACGGGGCCCCAGAGAAGGACGTGCTGGTGGCATCGGGCTGAGGCACCGCTCCCT
TTGGGACTACCAAGGCGCGTGCACCTGGCAGGACAAAGAGCATGAGTCTCGCTGGAG
ATCCAGGATCTGGCTGGAGGACTATGGGCTTACCGCTGTGAGGTATTGACGGCTGG
GGATGAAAGCGGCTGGTGGAGCTGGAGCTGCGGGGTGGTCTTCCCTTACAGTCCCCCA
ACGGCGCTACCACTTCAACTTCCACGAGGGCAGCAGGCTGTGCAAGAGCAGGCTGCC
GTGGCCTCTTGGAGCAGCTCTCCGGGCTGGAGGAGGGCTGGACTGGTGCAACGCC
CTGGCTGCAGGATGCTACGGTGCAGTACCCCATCATGGTCCCCGGCAGCCCTGCC
CAGGCCCTGCACCTGGCGTGCAGACTACGGCCCCGCCACCGCCGCTGCACCGCTATGAT
GTATTCTGCTTCGCTACTGCCCTCAAGGGGGGGTGTACTACCTGGACACCCCTGAGAAGCT
GACGCTGACAGAGGCAAGGGAGGGCTGCCAGGAAGATGATGCCACGATGCCAAGGTGGAC
AGCTCTTGGCGCTGGAGATTCCATGGCCTGGACCGCTGCACTGGCTGGCTGGAGAT
GGCAGCGTCCGCTACCTGTGGTTCACCCGATCTTAAGTGGGGCCCCAGAGCCTGGG
CCGAAGCTTGGCTTCCCCGACCCGAGAGCGCTGTACGGTTACTGCTACCGCCAGC
ACTAGGACCTGGGGCCCTCCCTGCCGATCCCTCACTGGCTGTGTATTATTGAGTGGTT
CGTTTCCCTTGTGGGGTGGAGCCATTAACTGTTTTATACTTCTCAATTAAATTCT
TTAACATTTTTACTATTTTGTAAAGCAAACAGAACCCAATGCCCTTGTCTCTG
GATGCCCACTCCAGGAATCATGCTTGTCCCCCTGGGCCATTGGGTTTGCTGGCTCTG
GAGGGTCTCCGCCATCAGGCTGGTCTCCCTCTTAAGGAGGTTGGTGGCCAGAGTGGC
GGTGGCTGTCTAGAATGCCGCCGGAGTCCGGCATGGGGCACAGTCTCCCTGCC
CAGCCTGGGGAGAACAGGGCCCTGGGGGCTCCGGAGCTGGCTTGGCCTCTCTGCC
CACCTCTACTCTGTGAAGCCGTGACCCAGTCTGCCACTGAGGGCTAGGGCTGGAA
GCCAGTTAGGCTTCCAGGCAGAACATCTGAGGGAGGAAGAAACTCCCTCCCGTTCCCT
TCCCTCTGGTTCAAAAGAACATCTGTTTGTCTATTGTTCTCTGTTCCCTGTGTGG
GGAGGGGCCCTCAGGTGTGTACTTGGACAATAATGGCTATGACTGCCCTCCGCCAA
AA
AA

FIGURE 78

MGLLLVPLLLLPGSYGLPFYNGFYYNSNSANDQNLGNHGKDLLNGVKLVVETPEETLFTYQ
GASVILPCRYRYEPALVSPRRVRVKWWKLSENGAPEKDVLVAIGLRHRSFGDYQGRVHLRQD
KEHDVSLEIQDLRLEDYGRYRCEVIDGLEDESGLVELELRGVVFYQSPNGRYQFNFHEGQQ
VCAEQAAVVASFEQLFRAWEGLDWCNAGWLQDATVQYPIMLPRQPCGGPGLAPGVRSGPR
HRLRHRYDVFCFATALKGRVYYLEHPEKLTITLEAREACQEDDATIAKVGQLFAAWKFHGLDR
CDAGWLADGSVRYPVVHPNCGPPPEPGVRSFGFPDPQSRLYGVYCQRQH

Signal sequence:

amino acids 1-17

Casein kinase II phosphorylation site.

amino acids 29-33, 53-57, 111-115, 278-282

Tyrosine kinase phosphorylation site.

amino acids 137-145

N-myristoylation site.

amino acids 36-42, 184-190, 208-214, 237-243, 297-303, 307-313

FIGURE 79

GGAGAGCGGAGCGAAGCTGGATAACAGGGGACCGATGATGTGGCGACCACATCAGTTCTGCTGC
TTCTGTTGCTACTGAGGCACGGGGCCCAAGGGAAAGCCATCCCCAGACGCAAGGCCCTCATGGC
CAGGGGAGGGTGCACCAGGCCCCCTGAGCGACGCTCCCCATGATGACGCCACGGAA
CTTCAGTAGCACCAGGCTTTCTGGGACGGGAAGTGGCAAGGAATTGACCAACTCA
CCCCAGAGGAAGGCCAGGCCCTCTGGGGGATCGTGACCGCATGGACCGCGCGGGGAC
GGCGACGGCTGGGTGTCGCTGGCCGAGCTTCGCGCGTGGATCGCGCACACGCAAGCGGCA
CATACGGGACTCGGTGAGCGGGCCTGGGACACGTACGACACGGACCGGACGGCGTGTGG
GTTGGGAGGAGCTCGCAACCCACCTATGGCACTACCGCCGGTGAAGAATTTCATGAC
GTGGAGGATCGAGACCTACAAAAGATGCTGGCTGGGACAGCGGGCTTCGGGTTG
CGACCAAGGATGGGACTCGATGGCACTCGAGAGGAGCTGACAGCCTTCCTGCACCCCGAGG
AGTTCCCTCACATGGGGACATCGTATTGCTGAACCCCTGGAGGACCTGGACAGAAACAAA
GATGGCTATGTCAGGTGGAGGAGTACATCGCGATCTGACTCAGCCGAGCCTGGGAGGA
GGAGCCGGCGTGGGTGCAAGACGGAGAGGAGCAGCTCCGGACTTCCGGATCTGAACAAGG
ATGGGCACCTGGATGGGAGTGAGGTGGGCACTGGGTGCTGCCCCCTGCCAGGACCA
CTGGTGGAAAGCCAACCACCTGTCACAGAGAGCGACAGACAAGGATGGCGCTGAGCAA
AGCGGAAATCTGGTAATTGGAACATGTTGTGGCAGTCAGGCCACCAACTATGGCGAGG
ACCTGACCCGGCACACGATGAGCTGTGAGCACCGCGCACCTGCCACAGCCTCAGAGGCC
CACAAATGACCGGAGGAGGGGGCGCTGGTCTGGCCCCCTCCCTGTCCAGGCCAGGAG
GCAGATGCACTCCAGGCATCCCTGGCTCTCAGGGACCCCTGGGTGCGCTTC
TGTCCCTGTCAACACCCCAACCCCAAGGGAGGGGCTGTCTAGTCCCAGAGGATAAGCAATAC
CTATTTCTGACTGAGTCTCCAGCCCAGACCCAGGGACCCCTGGCCCCAAGCTCAGCTCTAA
GAACCGCCCCAACCCCTCCAGCTCAAATCTGAGCCTCCACACATAGACTGAAACTCCCT
GGCCCCAGCCCTCTCTGGCTGGCTGGGACACCTCCCTCTGCCAGGAGGCAATAA
AAGCCAGCGGGGACCTTGAAAAAAAAAAAAAAAAAAAAAAA
AAAAAAAAAAAAAA

FIGURE 80

MMWRPSVLLLLLRLHGAQGKPSPDAGPHGQGRVHQAAPLSDAPHDDAHGNFQYDHEAFLGR
EVAKEFQDQLTPEESQARLGRIVDRMDRAGDGWVSLAELRAWIAHTQQRHIRDSVSAAWDT
YDTDRDGRVGWEELRNATYGHYAPGEEFHVEDAETYKKMLARDERRFRVADQGDGDSMATRE
ELTAFLHPEEFPHMRDIVIAETLEDLRNKGYVQVEEYIADLYSAEPGEEEPAWVQTERQQ
FRDFRDLNKDGHDGSEVGHVLPPAQDQPLVEANHLLHESDTDKDGRLSKAEILGNWNMFV
GSQATNYGEDLTRHHDEL

Signal sequence:

amino acids 1-20

N-glycosylation site.

amino acids 140-144

Casein kinase II phosphorylation site.

amino acids 72-76, 98-102, 127-131, 184-188, 208-212, 289-293,
291-295, 298-302

N-myristoylation site.

amino acids 263-269, 311-317

Endoplasmic reticulum targeting sequence.

amino acids 325-330

FIGURE 81

FIGURE 82

MSAAWIPALGLGVCLLLLPGPAGSEGAAPIAITCFTRGLDIRKEKADVLCPGGCPLLEFSVY
GNIVYASVSSICGAAVHRGVISNSGGPVRYSLPGRENYSSVDANGIQSQMLSRWSASFTVT
KGKSSTQEATGQAVSTAHPPTGKRLKKTPEKKTGNKDCKDIAFLIDGSFNIGQRRFNLQKN
FVGKVALMLGIGTEGPHVGLVQASEHPKIEFYLNKNTSAKDVLFIAIKEVGFRGGNSNTGKAL
KHTAQKFFTVDAGVRKGIPKVVVVFIDGWPSSDIEEAGIVAREFGVNVFIVSVAKPIPEELG
MVQDVTFVDKAVCRNNGFYSYHMPNWFGTTKYVKPLVQKLCTHEQMMCSTCYNSVNIAFLI
DGSSSGDSDNFRMLLEFVSNIAKTFEISDIGAKIAAVQFTYDQRTESFTDYSTKENVLAVI
RNIRYMSGGTATGDAISFTVRNVFGPIRESPNKNFLVIVTDQSYDDVQGPAAAHDAGITI
FSVGVAWAPLDDLKDMASKPKESHAFFTREFTGLEPIVSDVIRGICRDFLESQQ

Signal sequence:

amino acids 1-24

N-glycosylation site.

amino acids 100-104, 221-225

Casein kinase II phosphorylation site.

amino acids 102-106, 129-133, 224-228, 316-320, 377-381, 420-424,
425-429, 478-482, 528-532

N-myristoylation site.

amino acids 10-16, 23-29, 81-87, 135-141, 158-164, 205-211,
239-245, 240-246, 261-267, 403-409, 442-448, 443-449

Amidation site.

amino acids 145-149

FIGURE 83

CGCCGCGCTCCCGCACCGCGGCCCGCCACCGCGCCGTCCCGCATCTGCACCCGAGCCC
GGCGGCCCTCCCGCGGGAGCGAGCAGATCCAGTCGGGCCAGCGCACTCGGTCCAGTCG
GGCGCGCGCTCGGGCGCAGAGCGGAGATCGAGCGCTGGGCCACCCCTGCTGTGCGCTGC
TGCTGGCGCGCGGTCCCCACGGCCCCCGCCCGTCCAGCGGCACCTCGGCTCAGTC
AAGCCCGGCCCGCTCTCAGCTACCCCGAGGAGGAGGCCACCCCTCAATGAGATGTTCCCGGA
GGTTGAGGAAGCTGATGGAGGACACGCAGCACAATTGCGCAGCGCGTGGAAAGAGATGGAGG
CAGAGAAGCTGCTGCTAAAGCATCATCAGAAGTGAACTGGAAACTTACCTCCAGCTAT
CACAAATGAGAACACAGAACAGAAGCTGGACAAATGGTCTTTAGAGACAGTTACATCTG
TCACAAGATAACCAACACAGAACAGAAGCTGGACAAATGGTCTTTAGAGACAGTTACATCTG
TGGGAGACGAAAGGCAAGGGAGCAGAGCAGTGATCATCGACGAGGACTGTGGGCCAGC
ATGTAAGCCAGTTGCCAGCTTCCAGTACACCTGCCAGGCCATGCCGGGCCAGAGGATGCT
CTGCACCCGGGAGCAGTGCTGCTGGAGGCCAGCTGTGCTGGGGTCACTGCAACAAAAA
TGGCACCCAGGGGAGCAGAACGGGACCATCTGTGACAACACCAGGGACTGCCAGCGGGCTG
TGCTGCTGCCATGCCCGCCAGCCGGCTTCTGGACCTCATACCTGGGAGCTAGGCCATG
GCTTITGCCATGCCCGCCAGCCGGCTTCTGGACCTCATACCTGGGAGCTAGGCCATG
GAGCCTTGGACCATGGCTTCTGGCAGCTGGCTCTGCAACCCCCACAGCCACAGCTG
GTGTTATGTCAGACGCCACCTCTGGGGAGCCGCTGACCATGGGAGATGGGAGATCTG
CAGAGAGGTCCCCGATGAGTATGAAAGTGGCAGCTTCATGGAGGAGGTGCGCCAGGAGCTGG
AGGAACCTGGAGGGAGGCTGACTGAAGACATGGGAGCTGGGGAGCCCTGCGCTGCCCGCT
GCACGTGCTGGGGAGGGAGAGATTAGATCTGGACAGGCTGTGGTAGATGTCATAGAA
ATAGCTAATTATTCTCCAGGTGTCCTTAGCGTGGCTGACCAGGCTTCTCTAC
TCTCTTCCCAGTAAGTTCCCTCTGGCTTGACAGCATGGTGTGCACTTGTCTGC
TCCCTCCAGGTGTTCTCCAGGCTTCACAGTCTGGCTGGAGAGTCAGGCAGGGTTAAAC
TGCAAGGAGCAGTTGCCCCCTGTCAGATTATGGCTGCTTGCCTACCCAGTTGGCAG
ACAGCGTTGTTCTACATGGCTTGTAAATTGTTGAGGGAGGAGATGAAACAATG
AGTCTCCCTCTGATTGGTTTGGGAAATGTGGAGAGAGATGGCTTGGCAAACATCAA
CTGGCCTTGGGAAATGCAACAAATGAAATTCTCCAGCACTTCTCCATGGCATAGGT
TGCCCTCAGCTGTCAGATGAAATGTTCTGTCACCCCTGCATTACATGTTTATTCTAC
AGCAGTGTGCTCAGCTCTACCTCTGTGCGCAGGAGCATTCTCATATCCAAGATCAATT
CCTCTCTCAGCACAGCCTGGGGAGGGGTCTGGTCTCTCGTCATCAGGGATCTCAGAG
GCTCAGAGACTGCAAGCTGCTTGGCCAACTCACAGCTAGTGAAGACAGCAGCTTCT
CTGGTTGTGACTCTAACGCTCAGTGTCTCTCCACTAACCCACACCCAGCCTTGTGCCACCAA
AAGTGTCTCCCAAAAGGAGAAGGAGAATGGGATTCTTGAGGCATGCACATCTGGAAATTAAAG
GTCAAACATAATTCTCACATCCCTCTAAAGAAACTACTGTTAGGAACAGCAGTGTCTCAC
AGTGTGGGGAGCGCTCTCTAATGAGAACATGATATTGACACTGTCCTCTTGGCAGT
TGCATTAGTAATTGTAGGGCGAGGGATTAAATGAAATTGCAAATCACTTAGCAGAAC
GTACTTAGGTAAATTGTAGGGCGAGGGATTAAATGAAATTGCAAATCACTTAGCAGAAC
TGAAGACAATTATCAACCCAGCTGGAGAAAATCAAACCGAGCAGGGCTGTGTGAAACATGGTT
GTAATATGCGACTCGAACACTGAACTCTACGCCACTCCACAAATGATGTTTCAAGGTGTCA
TGGACTGTTGCCACCATGTATTCTACAGGTCTTAAAGTTAGTGTGACATGATTGTA
TAAGCATGCTTCTTGTAGTTAAATTATGATAAACATAAGTGTGCACTTGTGAAATCAAGC
ATAAAATCACTTCAACTGCAAAAAAAAAAAAAAAAAAAAAAA

FIGURE 84

MQRLGATLLCLLLAAAVPTAPAPAPTATSAPVKPGPALSYPQEEATLNEMFREVEELMEDTQ
HKLRSAVEEEMEAEEAAKASSEVNLPPSYHNETNTDKVGNNTIHVHREIHKITNNQTG
QMVFSETVITSGDEEGRRSHECIIDEDCGPSMYCQFASFQYTCQPCRGQRMLCTRSECCG
DQLCVWGHCTKMATRGSNGTICDNQRDCQPGLCFAQRGLLFPVCTPLPVEGELCHDPASRL
LDLITWELEPDGALDRCPCASGLLCQPHSHSLVYVCKPTFVGSRDQDGIELLPREVDEYEV
GSFMEEVVRQELEDLERSLTEEMALGEAAAAAALLGEEI

Signal sequence:

amino acids 1-19

N-glycosylation site.

amino acids 96-100, 106-110, 121-125, 204-208

Casein kinase II phosphorylation site.

amino acids 46-50, 67-71, 98-102, 135-139, 206-210, 312-316,
327-331

N-myristoylation site.

amino acids 202-208, 217-223

Amidation site.

amino acids 140-144

FIGURE 85

FIGURE 86

MRLLVAPLLLAWVAGATATVPVVPWHVPCPPQACQIRPWPYTPRSSYREATTVDCNDLFLTA
VPALPAGTQTLLQSNSIVRVDQSELGYLANLTLDLSQNSFSARDCDFHALPQLLSLHL
EENQLTRLEDHSFAGLASLQLBYLNHNQLYRIAPRAFGSLSNLLRLHLSNLLRAIDSRWFE
MLPNLEILMIGGNKVDAILDMNFRPLANLRSVLVAGMNLRREISDYALEGLQSLESLSFYDNO
LARVFPRRALEQVPGKLKFLDLNKNPLQRVGPQDFANMLHLKEGLNNNMEELVSIDKFALVNLP
ELTKLDITNNPLRSFIHPRAFPFHLPQMELTMMLNNNALSALHQQTVESLPNLQEVGLHGNPIR
CDCVIRWANATGTRVRFIEPQSTLCAEPPDLQRLPVREVPFREMTDHCLPLISPRSFPPSLQ
VASGESMVLHCRALAEPEPEIYWVTPAGLRLTPAHAGRRYRVPEGTLIELRVTAAEEAGLYT
CVAQNVLVGADTAKTVSVVVGRALLQPGDRQQGLELRVQETHYHILLSWVTPNTVSTNLTW
SSASSLRGQGATAALARLPRGTHSYNITRLQATEYWAQLOVAFADAHTQLACVWARTKEATS
CHRALGDRPGLIAILALAVLLAAGLA AHLGTGQPRKGVGRRPLPPAWAFWGWSAPSVRVV
SAPLVLPWNPGRKLPRSSEGETLPPQLSQNS

Signal sequence:

amino acids 1-18

Transmembrane domain:

amino acids 629-648

N-glycosylation site.

amino acids 94-98, 381-385, 555-559, 583-587

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 485-489

Casein kinase II phosphorylation site.

amino acids 46-50, 51-55, 96-100, 104-108, 130-134, 142-146,
243-247, 313-317, 488-492, 700-704

Tyrosine kinase phosphorylation site.

amino acids 532-540

N-myristoylation site.

amino acids 15-21, 493-499, 566-572

Amidation site.

amino acids 470-474, 660-664, 692-696

FIGURE 87

FIGURE 88

MRQTIIKVIKFILIICTVYVYVHNIFKFDVDCTVDIESLTGYRTYRCAHPLATLFKILASFYI
SLVIFYGLICMYTLWWMLRSSLKKYSFESIREESSYSDIPDVKNDFAFMLHLIDQYDPLYSK
RFAVFLSEVSENKLRLQLNLNNEWTLDKLQRQLTKNAQDKLELHLFMLSGIPDTVFDLVELEV
LKLELIPDVТИPPSIAQLTGLKELWLYHTAAKIEAPALAFRLRENLRALHIKFDTIKEIPLWI
YSLKTLEELHHTGNLSAENNRYIVIDGLRELKRKLVRLLKSNLSKLPQVVTDVGVHQLQKLSI
NNEGTKLIVLNLKMMANLTELIRCDLERPHSISFLSHNLQEIDLKDNNLKTIEEIISFQ
HLHRLTCLKLWYNHIAYIPIQIGNLTNERLYLNRNKEIEKIPTQLFYCRKLRYLDLSHNNLT
FLPADIGLQLNLQNLAITANRIETLPPPELFQCRKLRALHLGNNVLQSLPSRVGELETNLTQIE
LRGNRLECLPVELGECPLLKRSGLVVEEDLFNTLPPEVKERLWRADKEQA

Transmembrane domain:

amino acids 51-75 (type II)

N-glycosylation site.

amino acids 262-266, 290-294, 328-332, 396-400, 432-436, 491-495

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 85-89

Casein kinase II phosphorylation site.

amino acids 91-95, 97-101, 177-181, 253-257, 330-334, 364-368,
398-402, 493-497

N-myristoylation site.

amino acids 173-179, 261-267, 395-401, 441-447

FIGURE 89

GCCTGTTGCTGATGCTGCCGTGGTACTTGTCATGAGCTGGCACTGCCGCTCTCCCGT
CCCGCGGTGGTGTGCTGCCGTGGCTGAACGCAAGGAGCTGTCTATTGACT
GGCCCACAGAGGAGGGCAAGGAAGTATGGGATTATGTGACGGTCCGCAAGGATGCCATCATG
TTCTGGTGGCTTATTATGCCACCAACTCTGCAAGAACTTCAGAACTGCCCTGGTCAT
GTGGCTTCAGGGCGGTCCAGGGGTTCTAGCACTGGATTGGAAACTTGGAGGAAATTGGGC
CCCTTGACAGTGTACTCAAACCACGGAAAACCACCTGGCTCCAGGCTGCCAGTCTCTATT
GTGGATAATCCGTGGCACTGGGTTCACTTATGTGAATGGTAGTGGTGCCTATGCCAAGGA
CCTGGCTATGGTGGCTTCAGACATGATGGTTCTCTGAAGACCTTCTCAGTTGCCACAAAG
AATTCCAGACAGTCCATTCTACATTTCTCAGACTATGGAGGAAATGGCAGCTGGC
ATTGGTCTAGAGCTTATAAGGCCATTCA CGCAGGGACCATCAAGTGCACCTTGGGGGGT
TGCCCTGGGTGATTCTGGATCTCCCTGGATTGGTCTCTGGGACCTAACCTG
ACAGCATGTCCTTCTGAAGACAAAGGCTGGCAGAGGTGTCAGGTTGAGAGCAAGTA
CTGAATGCCGTAATAAGGGCTCTACAGAGGGCACAGAGCTGGGGAAAGCAGAAAT
GATCATTGAACAGAACACAGATGGGTGAACCTCTATAACATCTTAACAAAGCACTCCA
CGTCTACAATGGAGTCGAGTCTAGAATTCAACAGAGCCACCTAGTTGCTTTGTCAGGCG
CACGTGAGACACCTACAACGAGATGCCCTAACGGCAGCTCATGAATGCCCATCAGAAAGAA
GCTCAAAATTATTCTGGAGGATCAATCTGGGAGGCCAGGCTACCAACGCTTTGTAACAA
TGGAGGAGGACTTCATGAAGGCCAGTCATTGACATTGGACAGGTTGCTGGAGGCCAGGATC
AACGTGACCGTGTATAATGGACAGCTGGATCTCATCGTAGATACCATGGGTCAAGGAGCCTG
GGTGCAGAAACTGAAGTGGCCAGAACTGCCATTACAGTCAGCTGAAGTGGAAAGGCCCTGT
ACAGTGACCTAAATCTTGGAAACATCTGCTTTGTCAGTCAGTCTACAAAGACCTTGTCTTC
TACTGGATTCTGAAAGCTGGTCAATGGTCTCTGACCAAGGGACATGGCTCTGAAGAT
GATGAGACTGGTCACTCAGCAAGAATAGGATGGATGGGCTGGAGATGACCTGGTTGGCCT
TGGGGCACAGAGCTGAGCTGAGGCCGTGAAGCTGTAGGAAGGCCATTCTCCCTGTATCT
AACTGGGGCTGTGATCAAGAAGGTTCTGACCAAGCTCTGAGAGGATAAAATCATTGTCCT
GGAGGCAATTGGAAATTATTCTGCTCTTAAAAAAACCTAAGATTTTAAAAAATTGTAT
TTGTTTGATCAAATAAAGGATGATAATAGATATTAA

FIGURE 90

MELALRRSPVPRWLLLLPLLLGLNAGAVIDWPTEEGKEVWDYVTVRKDAYMFWWLYYATNSC
KNFSELPVWLQGGPGGSSTGFGNFEEIGPLDSLKPRTTLQAAASLLFDNPVGTFGSY
VNGSGAYAKDLAMVASDMMVLLKTFFSCHKEFQTVPFYIFSESYGGKMAAGIGLELYKAIQR
GTIKCNFAGVALGDSWISPVDVLSWGPYLYSMSLLEDKGLAEVSKVAEQVLNAVNKGLYRE
ATELGKAEMIIEQNTDGVNFYNIITKSTPTSTMESSLEFTQSHLVCLCQRHVRHLQRDALS
QLMNGPIRKKLKIIPEDQSWGGQATNVFVNMEEDFMKPVISIVDELLEAGINVTVYNGQLDL
IVDTMGQEAWVRKLWPELPKFSQLKWALKYSDPKSLETSAFVKSYKNLAFYWILKAGHMVP
SDQGDMALKMMRLLVTQQE

Signal sequence:

amino acids 1-25

N-glycosylation site.

amino acids 64-68, 126-130, 362-366

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 101-105

Casein kinase II phosphorylation site.

amino acids 204-208, 220-224, 280-284, 284-288, 351-355, 449-453

N-myristoylation site.

amino acids 22-28, 76-82, 79-85, 80-86, 119-125, 169-175,
187-193, 195-201, 331-337, 332-338, 360-366

FIGURE 91

GGCGCGGGAGAGGAGGCCATGGCGCGCGGGCGCTGCTGCTGGCCTGCTGGCT
GGGCTGGACTCAGGAAGCGGAGTCGCAGGAGGCGCGCCGTTATCAGGACCATGCGGCCGA
CGGGTCATCACGTGCGCATCGTGGGTGGAGAGGACGCCA**T**CAGGAGCCTGGCCGTGGCA
GGGGAGCCTGCGCCTGTTGGATTCCCACGTATGCGGAGTGAGCCTGCTCAGGCCACCGCTGGG
CACTCACGGCGCGCACTGCTTGAACCTATACTGACCTTAGTGTACCCCTGGGGTGGATG
GTCCAGTTGGCCAGCTGACTTCCATGCCATCCTCTGGAGCCTGCAGGCCACTACACCCG
TTACTTGTATCGAATATCTATCTGAGGCCCGCTACCTGGGAATTCAACCTATGACATTG
CCTTGGTGAAGCTGCTGCACTGTACACTAAACACATCCAGCCATCTGTCTCCAG
GCCCTCACATTGAGTTGAGAACCGGACAGACTGCTGGGTACTGGCTGGGGTACATCAA
AGAGGATGAGGCAC TGCCATCTCCCACACCCCTCCAGGAAGGACTCAGGTGCGCCATCAAACA
ACTCTATGTGCAACCCACCTCTCCTCAAGTACAGTTCCGAAGGACATCTTGGAGACATG
GTTTGCTGGCACGCCAACGGGAAAGGCGGGAGGATGCCCTGCTTGGTGACTCAGGTGGACCCCT
GGCCTGTAACAAGAATGGACTGTGGTATCAGATTGGAGTCGTGAGCTGGGAGTGGCTGTG
GTCGGCCAATCGGCCGGTGTCTACACCAATATCAGCCACCACCTTGAGTGGATCCAGAAG
CTGATGGCCAAGAGTGGCATGTCCAGCCAGCCCCCTCTGGCCACTACTCTTTTCCCTCT
TCTCTGGCTCTCCACTCCTGGGCCGGTTGAGCCTACCTGAGCCCATGCAGCCTGGGC
CACTGCCAAGTCAGGCCCTGGTCTTCTGTCTTGGTAATAAACACATTCCAGTTGA
TGCCCTTGAGGGCATTCTCAAAAAAAAAAAAAAAA

FIGURE 92

MGARGALLLALLLARAGLRKPESQEAAPLSGPCGRRVITSRIVGGEDAELGRWPWQGSRLW
DSHVCGVSSLSSHRWALTAACFETYSDLSDPSGMVQFGQLTSMPSFWSLQAYYTRYFVSNI
YLSPRYLGNSPYDIALVKLSAPVTYTKHIQPICLQASTFEFENRTDCWTGWGYIKEDEALP
SPHTLQEJVQVAIINNSMCNHLFLKYSFRKDIFGDMVCAGNAQGGKDACFGDSGGPLACNKNG
LWYQIGVVSVWGVGCGRPNRPGVYTNISHHEWIQKLMAQSGMSQPDPSWPLLFFPLLWALPL
LGPV

Signal sequence:

amino acids 1-18

N-glycosylation site.

amino acids 167-171, 200-204, 273-277

Casein kinase II phosphorylation site.

amino acids 86-90, 134-138, 161-165, 190-194, 291-295

N-myristoylation site.

amino acids 2-8, 44-50, 101-107, 225-231, 229-235, 239-245,
259-265, 269-275

Amidation site.

amino acids 33-37

Prokaryotic membrane lipoprotein lipid attachment site.

amino acids 252-263,

Serine proteases, trypsin family, histidine active site.

amino acids 78-84

FIGURE 93

CCCCACGGCTCCGCGGACGGTGGGAAGGGCAGAATGGACTCCAAGCTGCCCTCTAGGGCT
CTTTGCCCTCATCCTCTCGAAATGCGAGTTCAGCCGGAGGCCGACCGCGGAGGACG
TGCCCCCAGGCTGGGTGCCCCGGCGTCGGGAGCCTGAGGAAGAGCTGAGTCTCACCTT
GCCCTGAGACAGCAGAAATGTGGAAAAGACTCTCGGAGCTGGTGCAGGGTGTGCGGATCC
CTCTCCTCAATACGGAAAATACCTGACCCCTAGAGAATGTGCTGATCTGGTGAAGGCCATCCC
CACTGACCCCTCCACAGCGTCAAAGATGCTCTTGGCAGCGGAGGCCAGAACAGTCGATTCT
TGATCACACAGGACTTCTGACTGCTGGCTGAGCATCCGAGAACAGCTGCTGCTCC
TGGGGCTGAGTTCATCACTATGTGGGAGGACCTACGGAACCCATGTTGAAGTCCCCAC
ATCCCCTACAGCTTCCACAGGCTTGGCCCCCATGTGGACTTGTGGGGGACTGCACCGT
TTTCCCCAACATCATCTGGCAAGCTCTGGCAGGGTGCAGGTGACAGGGACTGTAGGCC
GCATCTGGGGAACCCCTCTGATCCGTAAGGATACAACTTGACCTCAAGAGCTGG
GCTCTGGCACCAAGCAATAACAGCCAAGCTGTGCCAGTCTCTGGAGCAGTATTTCCATGAC
TCAGACCTGGCTGAGTTCATGCCCTCTGGCAACTTGGCAGCATCAGGATCAGTAC
CCGGTGGTGGTGGACAACAGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG
ACCTGATGAGTGTCTGGCCAACATCTCCACCTGGTCTACAGTAGGCCCCGGCATGAG
GGACAGGGGCCCTCTCTGCACTGGCTCACTGCTCTGAGTAATGAGTCAGGCCCTGG
GCATACTCTGGAGCTATGGAGATGAGGACTCTCTGGCAGCAGGCCATCATCAGGGGTC
ACACTGGAGCTCATGGGCTGGCGCTGGGGTCTACCCCTGCTCTGGCTCAGGGAGCAGT
GGGGCCGGGTGGTGGTCTCTGGAGACACAGCTCCGGCTACCTCCCTGCTCC
CCCTATGTCAACAGCTGGAGGACATCCCTGGAGAACACTTCTCCATCACAATGAAA
TTGGTGAATATATCAGTGGTGGTGGCTCAGGAATGTTCCACGGCCTTCATACAGGAG
GAAGCTGTAACGAAGGTTCTGAGCTCTAGCCCCCACCTGGCCACATCAGGTTACT
CAATGCACTGGCCCTGCTACCCAGATGTGGCTGCACTTCTGATGGCTACTGGTGGTCA
GAGTGGCCATTCTGGGTTGGGCTGGGAACTCCGGCTCTACTCTCAGTTGGGGGATCTTA
TCTCTGATCATGAGCACAGGATCTTGGCCGGGGCTTCTGGGTTGGCTTCAACCGA
GCTCTACCCAGCAGCATGGGGCAGGTCTTGTGATGTAACCCGTGGCTGCATGAGTCTG
TGGATGAAAGGGTAGAGGGGCCAGGGTTCTGCTCTGGTCTGGCTGGGATCTGTAACAGG
TGGGGAAACCCAATCTCCAGCTTGTGAGACTCTACTCAACCCCTGACCCCTTCTCATC
AGAGGAGATGGCTTGTCTGGCTCTGGCTGAACCTCAACTTGTGCTGAGCAGCTT
GAAGGCCCTGCTGAACCCCTCAACTTGTGACTGCTGAGACAGCTTATCTCCCTAACCC
TGGCTGTGAGCTTGTGACTCCAAACCTTGTGCTCCATCATACTCAGGCTCTCC
CTCTGCTTGAATCTTGTCAATAGATGCTGTAACTAGCATTTTGTGAATGCTCT
ATCTCATCTTCTCTTCAATCAGGCTTTCCAAGGGTTGATACAGACTCTGTGACTA
TTTCACTGATATTCACTCCCAATTCACTGCAAGGGACCTACTGTCACCGTTACTCT
TTCTTACCCCTGACATCCGAAACATGGCTTCTGGCTTCACTGTCATCTCAATCTT
GCCCTTCCATCATGATGGCCCACTCTCTCTACTTAGCTTCCAGGCTTAACTCTCTG
ACTACTCTGTCTTCTCTCTCATCAATTCTGCTTCTCATGGAATGCTGACCTTATTG
TCCATTGATATTGCTCTTCTCACTGTTACTCATGTCCTGGACAAATCAGTGC
TCTACAAACATTACCATCTCACTAAAGACTTTCTATCCAAATAATGATTGATACTCT
TGTAAAAAA

FIGURE 94

MGLQACLLGLFALILSGKCSYSPEPDQRRTLPPGVSLGRADPEEEELSTFALRQQNVERLS
ELVQAVSDPSPQYGKYLTLENVAIDLVRSPSPLTLHHTVQKWLLAAGAQKCHSVITQDFLTCWL
SIRQAELLPLGAEFHYYVGGPTEHVVRSPHPYQLPQALAPHVDFVGGHLRFPPPTSSLRQRP
EPQVTGTVGLHGVTPSVIRKRYNLTSQDVSGTSNNSQACAQFLEQYFHDSDLAQFMRLFG
GNFAHQASVARVVGQQGRGRAGIEASLDVQYLMSAGANISTWVYSSPGRHEGQEFLQWLML
LSNESALPHVHTVSYGDDEDSLSAYIQRVNTELMAAARGLTLFASGDGAGCWSVSGRH
QFRPTFPASSPYVTTVGGTSFQEPFLITNEIVDYISGGGFSNVFPRPSYQEEAVTKFLSSSP
HLPPSSYFNASGRAYPDVAALSDGYWVVSNRVPIPWVSGTSASTPVFGGILSLINEHRILSG
RPPLGFLNPRLYQQHGAGLFDVTRGCHESCLEEVVEGQQFCSGPGWDPVTGWGTPTSQCL

Signal sequence:

amino acids 1-16

N-glycosylation site.

amino acids 210-214, 222-226, 286-290, 313-317, 443-447

Glycosaminoglycan attachment site.

amino acids 361-365, 408-412, 538-542

Casein kinase II phosphorylation site.

amino acids 212-216, 324-328, 392-396, 420-424, 525-529

N-myristoylation site.

amino acids 2-8, 107-113, 195-201, 199-205, 217-223, 219-225,
248-254, 270-276, 284-290, 409-415, 410-416, 473-479, 482-488,
521-527, 533-539, 549-555

FIGURE 95

GGCGCGCGCTCTCTCCGGGCCACACCTGTCGAGCGCGCAGCGAGCCGCCGGGG
GGGCTGCTCGGCGCGAACAGTGCTCGGAATGGCAGGGATTCCAGGGCTCTTCTCT
TTCTTTCTGCTCTGTGCTGTTGGCAAGTGAGCCCTACAGTCCCCCTGGAAACCCACTTG
GCCCTGCATAACCCTCCCTGCTGCTTGGCCAGTCAACCTCAATTAGCCAAGCCAGACT
TTGGAGCCGAAGCCAATTAGAAGTATCTCTCATGTTGGACCCCAGTGTCTATGCCAATGG
CCACTGCCCACCTACGAAGAGGCCAAGCAATATCTGCTTATGAAACGCTCTATGCCAATGG
CAGCCGCACAGAGACGCAAGGTGGGCATCACATCCTCAGCAGTAGTGGAGATGGGCCAAC
ACCGAGACTCAGGGTCTCAGGAAAGTCTCGAAGGAAGCGGCAGATTATGGCTATGACAGC
AGGTTCACTTGGGAAGGACTTCTGCTCAACTACCCCTTCTAACATCAGTGAAGTT
ATCCACGGGCTGCACCGCACCTGGTGGCAGAGAACATGTCCTCACAGCTGCCACTGCA
TACACGATGGAAAACCTATGTGAAAGGAACCCAGAAGCTTCAGTAGTGGCTTCCCTAAAGGCC
AAGTTAAAGATGGTGGCAGGGGCCAACGACTCCACTTGCCATGCCAGAGCATGAA
ATTTCACTGGATCCGGGTGAAACGACCCATGTGCCAAGGGTGGATCAAGGGCAATGCCA
ATGACATCGCATGGATTATGATTATGCCCTCTGGAACTCAAAAGCCCCACAAGAGAAA
TTTATGAAGATTGGGGTGAGCCCTCTGCTAAGCAGTGCCAGGGGCCAGAATTCACTTCTC
TGGTTATGACAATGACCGACCAAGCAATTGGTGTATCGCTCTGTCAGCTCAAAGACGAGA
CCTATGACTGCTCTACCAAGCAATGCGATGCCAGCCAGGGCCAGGGCTGGGTCTAT
GTGAGGATGTGGAAGAGACAGCAGCAGAAGTGGGAGGCCAAAATTATTGGCATTTTTCA
GCACCACTGGGTGGACATGAATGGTCCCCACAGGATTCAACGTGGCTGTCAAAATCACTC
CTCTCAAATATGCCAGATTGCTATTGGATTAAAGGAAACTACCTGGATTGTAGGGAGGG
TGACACAGTGTCCCTCTGGCAGCAATTAGGGCTTCATGTTCTTATTTAGGAGAGGCC
AAATTGTTTTGTCTTGGCTGCACAGTGTGTGTGTGTGTGTGTGTAAAGGTG
CTTATAATTTTACCTTTCTACATTGCAAGATGACTGGTTACTATTGAAAAGT
GTTTGTGTATCATATCATATCATTTAAGCAGTTGAGGCATACTTTGCTAGAAATAA
AAAAAAACTGATTGGGCAATGAGGAATATTGCAAACTTAAGTAAATTCTCAGGTTTG
CAAACTTGATTTTATTCATGCAACTGTTCAAAGATTATTAATATTGCA
CAAGAGATGAAAAAAAAAAAAAA

FIGURE 96

MAGIPGLLFLFFFLLCAVGQVSPYSAPWKPTWPAYRLPVVLQPSTLNLA
KPDFGAEAKLEVS
SSCGPQCHKGTPPLPTYEEAKQYLSYETLYANGSRTETQVGIYILSSSGDGAQHRD
SGSSGKS
RRKRQIYGYDSRFSIFGKDPLLNPFS
TQKLRVGFLPKFKDGGRGANDSTS
SAMPEQMKFQWIRVKRTHVPKGWIKGNANDIGMDYDYA
LLELKPKH
AQPGASGSGVYV
IKGNYLD
RMWKRQQQKWERKIIGIFSGHQV
DLYQQCD
VDMNGSPQDFNV
AVRITPLKYAQICYW
REG

Signal sequence:

amino acids 1-19

N-glycosylation site.

amino acids 93-97, 207-211

Glycosaminoglycan attachment site.

amino acids 109-113, 316-320

Casein kinase II phosphorylation site.

amino acids 77-81, 95-99, 108-112, 280-284, 351-355

N-myristoylation site.

amino acids 159-165, 162-168, 202-208, 205-211, 314-320, 338-344

Serine proteases, trypsin family, histidine active site.

amino acids 171-177

FIGURE 97

GCATCGCCCTGGGCTCTCGAGCCTGCTGCCGTCCCCCGCCCCACAGCCATGGGGTT
CTGGAGCGCCCCCAGCCCTGGGTGGGGCTGTCGGCACCTTCACCTCCCTGCTGCTGCTG
GCCTCGACAGCCATCCTCAATCGGGCAGGATACTCTGTCCTCCCAGGCTGTGGGAAGCCCCA
GCAGCTGAACGGGTTGGGGGGCAGGACAGCACTGACAGCGAGTGGCCCTGGATCGTGA
GCATCCAGAAGAAATGGGACCCACCCTGCGAGGTTCTGCTCACAGCCGCTGGGTGATC
ACTGCTGCCACTGTTCAAGGACAACCTGAACAAACCATACCTGTCCTGTGCTGCTGGG
GGCCTGGCAGCTGGGAAACCTGGCTCTGGTCCCAGAAGGTGGGTGGCTGGGAGC
CCCACCCCTGTATTCTTGAAGGAAGGTGCTGTGAGACATTGCCCTGGTGCGTCTCGAG
CGCTCCATACAGTTCTCAGAGCGGGTCTGCCCATCTGCCACTCTGATGCCCTATCCACCT
CCCTCAAACACCCACTGTTGGATCTCAGGCTGGGGAGCATCCAAGATGGAGTCCCTTG
CCCACCCCTAGACCCCTGAGAACGCTGAAGGTTCTATCATGACTCGGAAGTCTGAGCCAT
CTGTACTGGGGGAGCAGGACAGGGACCCATCACTGAGGACATGCTGTGCGCGCTACTT
GGAGGGGGAGGGGATGCTTGTCTGGGACTCGGGGCCCTCATGTCGCCAGGGACG
GCGCTGGCTGGCGCATCATCAGCTGGGGAGGGCTGTGCCGAGCGAACAGGCC
GGGGTCTACATCAGCCTCTGGCACCGCTCTGGGTGAGAACATCGTCAAGGGGTGCA
GCTCCGCGGGCGCTCAGGGGGTGGGCCCTCAGGGCACCGAGCCAGGGCTGGGCCG
CCGCGCGCTCTAGGGCGCAGCGGGACGCCAGGGCTGGATCTGAAAGGCGCCAGATCCACA
TCTGGATCTGGATCTGGCGGGCTCGGGCGTTCCCCCGCGTAAATAGGCTCATCTACC
TCTACCTCTGGGGCCCGACGGCTGCTGGAAAGGAACCCCTCCCGACCCGCCGAC
GGCCCTCAGGCCCCCTCCAAGGCATCAGGCCCCGCCAACGGCCTCATGTCCTGCC
GACTTCCGGCCCCGGGGCCCCAGCGCTTTGTGTATATAATGTTAATGATTTTAT
AGGTATTGTAACCCCTGCCACATATCTTATTCTTCTCCAATTCAATAAAATTATTTATT
CTCCAAAAAAA

FIGURE 98

```
></usr/seqdb2/sst/DNA/Dnaseqs.full/ss.DNA43318
><subunit 1 of 1, 317 aa, 1 stop
><MW: 33732, pI: 7.90, NX(S/T): 1
MVGSGAPPALGGGCLGTFTSLLLLASTAILNAARIPVPPACGKPQQLNRVVGGEDSTDSEWP
WIVSIQKNGTHHCAGSLLTSRWVITAHHCFKDNLNKPYLFSVLLGAWQLGNPGSRQKVGVVA
WVEPHPVYSWKEGACADIALVRLERSIQFSERVLPICLPDASIHLPPNTHCWISGWGSIQDG
VPLPHPQTLQKLKVPIIDSEVCSHLYWRGAGQGPITEDMLCAGYLEGERDACLGDSGGPLMC
QVDGAWLLAGIISWGEGCAERNRPGVYISLSAHRSWVEKIVQGVQLRGRAQGGGALARPSQG
SGAAARS
```

Signal sequence:

amino acids 1-32

N-glycosylation site.

amino acids 62-66, 96-100, 214-218, 382-386, 409-413, 455-459,
628-632, 669-673, 845-849, 927-931, 939-943, 956-960

Glycosaminoglycan attachment site.

amino acids 826-830

Casein kinase II phosphorylation site.

amino acids 17-21, 39-43, 120-124, 203-207, 254-258, 264-268,
314-318, 323-327, 347-351, 464-468, 548-552, 632-636, 649-653,
671-675, 739-743, 783-787, 803-807, 847-851, 943-947, 958-962,
1013-1017, 1019-1023, 1021-1025

Tyrosine kinase phosphorylation site.

amino acids 607-615

N-myristoylation site.

amino acids 179-185, 197-203, 320-326, 367-373, 453-459, 528-534,
612-618, 623-629, 714-720, 873-879

FIGURE 99

GACGGCTGCCACCATGCACGGCTCCTGCAGTTCCCTGATGCTTCTGCTGCCGCTACTGCTA
CTGCTGGTGGCCACCACAGGCCCGTGGAGCCCTCACAGATGAGGAGAACGTTGATGGT
GGAGCTGCACAAACCTCTACCGGGCCCAGGTATCCCGACGGCCTCAGACATGCTGCACATGA
GATGGGACGAGGAGCTGGCCGCCCTCGCCAAGGCCTACGCACGGCAGTGCCTGTGGGCCAC
AACAGGAGCGGGCGGCCGGCGAGAATCTGTCGCCATCACAGACGAGGGCATGGACGT
GCCGCTGGCCATGGAGGAGTGGCACACAGCGCTGAGCACTACAACCTCAGCGCCGCCACCT
GCAGCCAGGCCAGATGTGCGGCCACTACACGCAGGTGGTATGGCCAAGACAGAGAGGATC
GGCTGTGTTCCACCTCTGTGAGAAGCTCAGGGTGTGAGGAGACCAACATGAAATTACT
GGTGTGCAACTATGAGCCTCCGGGAACGTAAGGGAAACGGCCCTACCGAGGGGACTC
CGTGCTCCCAATGTCCTCTGGCTACCACGTCAAGAACCTCCCTGTGAACCCATCGAACG
CCGGAAGATGCTCAGGATTGCTTAACCTGTAACGTAGGCCCATCCTCCGGCGACTGA
AGCATCAGACTCTAGGAAATGGTACTCCTCTCCCTAGCAACGGGATTCGGCTTCT
TGGTAACAGAGGTCTCAGGCTCCCTGGCAACCAAGGCTCGCCTGTGAAACCCAGGCC
CCAACCTCCCTAGCAACGAAAGACCCGCCCTCCATGGCAACAGAGGCTCCACCTTGCGTAAC
AACTGAGGTCCCTCCATTGGCAGCTCACAGCCTGCCCTGGATGAGGAGCCAGTTA
CCTTCCCCAAATGACCCATGTCCTATCCAAAATCAGCAGACAAAGTGACAGACAAAAACA
AAAGTGCCTCTAGGAGGCCAGAGAACTCTGACCCCCAAGATGTCCCTGACAGGGCAAG
GGAACCTCCTACCCCATGCCAGGAGGAGGTGAGGCTGAGGCTGAGTTGCCCTTCAGTG
AGGTCTTGGCCTCAGTTTCCAGGCCAGGACAAGCCAGGTGAGCTGCAGGCCACACTGGAC
CACACGGGGCACACCTCCTCCAAGTCCCTGCCAATTCCCAAATACCTCTGCCACCGCTAA
TGCCACGGTGGCGTGCCTGGCTCGCAGTCCTTGCCAGGTGCAAGAGGCCCTGACA
AGCCTAGCGTTGTGTCAGGGCTGAACCTGGCCCTGGCATGTGTGGGCCCTCTCTGGGA
CTACTGCTCCTGCCTCCTCTGGTGTGGCTGGAATCTTCTGAATGGGATACCAACTCAAAGGG
TGAAGAGGTCACTGCTCCTCTGTCACTTCCCCACCTGCCCCAGCCCTAAACAAGATA
CTTCTTGGTTAAGGCCCTCCGAAGGGAAAGGCTACGGGCATGTGCCCTCATCACACCATCC
ATCCTGGAGGCACAAGGCCCTGGCTGGCTGGAGCTCAGGAGGCCCTGAGGACTGCCACACC
GGGCCACACCTCTCCTGCCCTCCCTGGAGTCCTGGGGGTGGAGGAGTTGAGGGAGCT
CACTGCCCTACCTGGCCTGGGCTGTCTGCCACACAGCATGTGCCTCTCCCTGAGTGCCTG
TGTAGCTGGGATGGGATTCCCTAGGGCAGATGAAGGACAAGCCCCACTGGAGTGGGTT
TTTGAGTGGGGAGGCAGGGAGGAAAGTAACCTGACTCTCCAATAAAACCT
GTCCAACCTGTGAAA

FIGURE 100

MHGCSFLMLLLPLLLLWATTGPVGALTDEEKRLMVELHNLYRAQVSPTASDMLHMRWDEE
LAAFAKAYARQCVCWGHNKERGRRGENLFAITDEGMDVPLAMEEWHEHEREHYNLSAATCSPGQ
MCGHYTQVVWAKTERIGCGSHFCEKLQGVETNIELLVCNYEPPGNVKGKRPyQEGTPCSQC
PSGYHCKNSLCPIGSPEADAQDPYLVTEAPSFRATEASDSRKMGTPSSLATGIPAFLVTEV
SGSLATKALPAVETQAPTSLATKDPPSMATEAPPCTEVPSILAAHSIPLSDEEPVTTPKS
THVPIPKSADKVTDKTKVPSRSPENSLDPKMSLTGARELLPHAQEEAEAEELPPSSEVLAS
VFPAQDKPGELQATLDHTGHTSSKSLPNFPNTSATANATGGRALALQSSLPGAEGPDKPSVV
SGLNNSGPGHVWGPILLGLLLLPLVLAGIF

Signal sequence:

amino acids 1-22

N-glycosylation site.

amino acids 114-118, 403-407, 409-413

Glycosaminoglycan attachment site.

amino acids 439-443

Casein kinase II phosphorylation site.

amino acids 29-33, 50-54, 156-160, 195-199, 202-206, 299-303

N-myristoylation site.

amino acids 123-129, 143-149, 152-158, 169-175, 180-186, 231-237,
250-256

Amidation site.

amino acids 82-86, 172-176

Peroxidases proximal heme-ligand signature.

amino acids 287-298

Extracellular proteins SCP/Tpx-1/Ag5/PR-1/Sc7 signature 1.

amino acids 127-138

Extracellular proteins SCP/Tpx-1/Ag5/PR-1/Sc7 signature 2.

amino acids 160-172

FIGURE 101

FIGURE 102

MVDVLLLFLSCLLFHISRPDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVSAN
ITLSSLAGNRIVEILPEHLKEFQSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVTSMEP
GYFDNLANTLLVVKLNRRNISAIPPKMFKLPQLQHLELNRNKIKNVGDLTFQGLGALKSLKM
QRNGVTKLMGAFWGLSNMEILQLDHNLTEITKGWLGYGMLQELHLSQNAINRISPDAWE
FCQKLSELDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRCGLSSLKTLDDLKNNE
ISWTIEDMNGAFSGLDKLRRLLIQGNRIRSIITKKAFTGLDALEHLDLSDNAIMSLQGNAFSQ
MKKLQQLHLNTSLLCDCQLKWLQPWVAENNQFSFVNACAHQPLLKGRSIFAVSPDGTVCD
DFFPKPQITVQPETQSAIKGSNLSCICSAASSSDSPMTFAWKDNELLHDAEMENYAHLRAQG
GEVMEYTTILRLREVEFASEGKYQCVISNHFGSSYSVKAKLTVNMLPSFTKTPMDLTIRAGA
MARLECAAVGHFAPQIAWQKDGGTDFPAARERRHMHVMPEDDVFFFIVDVKIEDIGVYSCAQN
SAGSISANATLTVLETPSFRLPULLDRTVTKGETAVLQCIAGGSPPPKLNWTKDDSPLVUTER
HFFAAGNQLLIIVDSDVSDAGKYTCMSNTLTERGVNRLSVIPTPTCDSPQMNTAPSLLDDG
WATVGVVIIAVVCCVVGTSLVVVVIIYHTRRRNEDCSITNDTETNLPADIPSYLSSQGTLLAD
RQDGYVSSESQSHHQFVTSSGAGFFLPQHDSGGTCHIDNSSEADVEATDLFLCPFLGSTGP
MYLKGNVYGSDFETYHTGCSPPDRTVLMDHYEPSYIKKECYPCSVHSPEESERSFSNISW
PSHVRLKLLNTSYSHNEGPGMKNLCLNKSSLDFSANPEPASVASSNSFMCTFGKALRRPHLDA
YSSFGQPSDCQPRAFYLAHSSPDLDSGSEEDGKERTDFQEENHICTFKQTLNEYRTPNFQS
YDLDLT

Signal sequence:

amino acids 1-19

Transmembrane domain:

amino acids 746-765

N-glycosylation site.

amino acids 62-66, 96-100, 214-220, 382-386, 409-413, 455-459,
628-632, 669-673, 845-849, 927-931, 939-943, 956-960

Glycosaminoglycan attachment site.

amino acids 826-830

Casein kinase II phosphorylation site.

amino acids 17-21, 39-43, 120-124, 203-207, 254-258, 264-268,
314-318, 323-327, 347-351, 464-468, 548-552, 632-636, 649-653,
671-675, 739-743, 783-787, 803-807, 847-851, 943-947, 958-962,
1013-1017, 1019-1023, 1021-1025

Tyrosine kinase phosphorylation site.

amino acids 607-615

N-myristoylation site.

amino acids 179-185, 197-203, 320-326, 367-373, 453-459, 528-534,
612-618, 623-629, 714-720, 873-879

FIGURE 103

GGGGAGAGGAATTGACCATGTAAAAGGGACTTTTCTTGGTGGTGGCTGTTGGTGCTTCGCAAAATG
AAGGATGCGAGGCAGCTTCTCTGGAACCGAACGAAATGATAAATGATTGTGCAAGAGAGAAGGAGAAC
GAAGCTTCTTCTTGTGAGCCCTGGATCTAACACAAATGTGTATATGTGCAACAGGGAGCATTCAGAAGTAAA
TAACCAGAGTGTAGACCCGGGGGTGGTGTCTGACAATAAATATCTTAAAGCAGCTGTCCTCCCTC
CCCCCCCCAAAAAAAGGATGATGGAAATGAAAGAACCGAGGATTCAAAAGAAAAAATGTATGTCATTCTC
TATAAGGAGAAGTGGAGGAGATTTTGGATGAAAAGTTGGGCTTTTGTAAAGTAAAGTAAGAACT
GGTGTGGTGTGTTCTTCTTCTTGAATTTCCCACAGGAGAGGAAATTAATAATACATCTGCAAGAAA
TTTCAGAGAAGAAGTGTGACCGGGCAGATGGAGCATTGAGGATTTCTTCATCACCTCTTAAAT
TTTGTGCCATATGTGACTAAATGAGGATAATGCGATGTTGATTTCTTCATCACCTCTTAAAT
TTTATCCCTTGTGATCAAGATCATGGTTCTTGTCTTAAACACCTGGATCTTCATCTGGATGTGCT
GTGATCAGTAAATACAATCTGGTAAAGGACCAACAGGAGATAAATTATGAAATGTTGAAAGAAGAT
GACCTTACATCCACAGCAGATAATGAGTCTGGGCTAGTTAACAGGGCCCTTTGACCCCTGCTGTGCT
GCTGGCTCTCAACTTCTGTGGTGTGGCTGGTCTGGGCTGGGCTCTCTGTGCTCTGAGCAA
CCAGTTCAGCAGGATGGTGTGGTGTGGAAACCTCGCTGGATGGCATCTCCACCAACACGGCT
GCTGAACCTCATGAGAACCAAATCAGATCATCAAAGTGAACAGCTCAAGCAGTGGGACTTGGAAATCT
ACAGTTGAGTAGGAAACCTATCAGAACCTTGAATTTGGGCTTCAAGTGTCTGGCAACACTCAACACTCTGGA
ACTCTTGAATCTTCTTACTACATCCGAGATGAGGATTTGTATCTGCTTAAACTGAGGAGCTCTGGT
GGAAACAAACCCCTGGAAAGCATCCTTCTTATGCTTTAACAGAAATCCCTCTTGGCCGACTAGACCTGG
GGAATGAAAAGACTCTCATCATCTGCAAGAGGCTCTTGAAGGCTCTGGATCACTTGAAGGTTTGAACCTTG
CATGTCGAACCTTGGGAAATCCCTAACCTCACCCGCTCATAAAATAGATGAGGTGGATCTTCTGGGAATCA
TTTATCTGCCATCAGGGCTGGCTGGGGTGTGGATGACCTTCAAAACTGGGATGATGATACAGTCCAGAT
TCAGTGTGATGAAAGCCTTCTGACAACCTTCACTGGGACTGAGGATGATCAACTGGCACACAATAATCAAC
ATTACTGCTCATGACCTTCTACTCCCTGATCATCTGGGAGGACATACATGGGAGGACATACATTACATGCTGG
TACACTCTGACATCTGGCTCAGCTGGTGGGATAAAAGACATGGCCCCCTGAACACAGCTTGTGCCCCGGT
TACACTCTCCCATCTAAAGGGAGGTACATTGGAGAGCTGACAGAAATTACTTCACATGCTATGCTCCGGT
GATTTGGAGCCCCCTGCAAGACCTCAATGTCAGGAGCTGACAGGAGCTGGAGCTGAGCTGAAATGTGCGGCTC
CTTGACATCTGATCTGGATACTCTGGAAATGCAAGACCTGAGCTGACACATGGGGCTGACAAAGTGCAGGATGCTG
GCTCAGTGTGATGGTACGTTAAATTCACAAATGTAACTGTGCAAGGATACAGGCTGACATGACACATGGGGCTGACAAAGTGC
TTCCGGTGGGAATACTACTGCTTCAGGCCACCTGGAATGTTACTCTGGACACCTACTCTTCTTACTTTT
AACCGTCACAGTAGAGACTGGAACCGTCTCAGGATGAGGACAGGACCAAGATAACATGTGGGCTCCACTCC
AGTGGCAGACTGGGAGACCAACATGGAACCGTCTCAGGATGAGGACAGGACCAAGATAACATGTGGGCTCCACTCC
CACCATCCCACTGACTGATATAAACATGGGACCTGGGAAATGTGAGGATCATGAGACTACCAAAATCATCAT
TGGGTGTTCTGGCCATCACACTCATGGCTGCACTGATGCTGGTCACTTACAAAGATGAGGAAAGCAGCAGCA
TCGGAAAACCATCACGGCCACAAAGGACTGTGAAATATTATAATGTGAGGATGAGGATACGGGAGACACACC
CATGGAAAGGCCACTGGCCATGCTGCTATGGAGGATGAGGACCTAAATCACTATAACTCATACAAATCTCCCT
CAACACACAAACAGTTAACACATAAAATCAATACAGCTGAGCTGAGGATGAGGATACGGGTTATGATCCGAATGAA
CTCTAAAGACAATGACAGAGACTCAAATCTAAACATTACAGAGTTACAAAAAAACAAACATCAAAAAAA
GACAGTTATTTAAATGACACAAATGACTGGGCTAAATCTACTGTTCAAAAAAAGTGTCTTACAAAAAA
AAAAAGAAAAGAAAATTTTTATTAAAAAAATTCTATGTGATCTAAAGCAGACAAAAAA

FIGURE 104

MLNKMTLHPQQIMIGPRFNRALFDPLLVLLALQLLVAGLVRAGTCPSVCSNQFSKVIC
VRKNLREVPDGISTNTRLLNLHENQIQIKVNSFKHLRHEILQLSRNHIRTIEIGAFNGLA
NLNTLELFDNRLTTIPNGAFVYLSKLKELWLRNNPIESIPSYAFNRIPSLRRLDLGELKRLS
YISEGAFEGLSNLRYLNLMCNLREIPNLTPLIKLDELDLSGNHLSAIRPGSFQGLMHLQKL
WMIQSQIQVIERNAFDNLQSLVEINLAHNNTLLPHDLFTPPLHHLERIHLHHNPWNNCNDIL
WLSWWIKDMAPSNTACCACRNTPPNLKGRYIGEIQDQNYFTCYAPVIVEPPADLNVTTEGMAAE
LKCRASTSLTSVSWITPNGTVMTHGAYKVRIAVLSDGTLNFTNVTVQDTGMYTCMVNSVGN
TTASATLNVTAAATTTFSYFSTVTVETMEPSQDEARTTDNNVGPTPVVDWETTNVITSLTPQ
STRSTEKTFTIPTVDINSGIPGIDEVMKTTKIIIGCFVAITLMAAVMLVIFYKMRKQHHRQN
HHAPTRTVEIINVDDIEITGDTPMESHLPMPAIEHEHLNHYNSYKSPFNHTTVNTINSIHSS
VHEPLLIRMNSKDGVQETQI

Signal sequence:

amino acids 1-44

Transmembrane domain:

amino acids 523-543

N-glycosylation site.

amino acids 278-282, 364-368, 390-394, 412-416, 415-419, 434-438,
442-446, 488-492, 606-610

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 183-187

Casein kinase II phosphorylation site.

amino acids 268-272, 417-421, 465-469, 579-583, 620-624

N-myristoylation site.

amino acids 40-46, 73-79, 118-124, 191-197, 228-234, 237-243,
391-397, 422-428, 433-439, 531-537

FIGURE 105

FIGURE 106

MASAPSLRARAAAGLGLLLCAVLGRAGRSDSGGRGELCQPSGVAEARPCPTTCRCLGDLDDCSR
KRLARLPEPLPSWVARLDLSHNRNLSFIKASSMSHLQSLEVKLNNELETIPLNLGPVSANIT
LLSLAGNRIVEILPEHLKEFQSLETLDLSSNNISELQTAFFPALQLKLYLNSNRVTSMEPGY
FDNLANTLLVLKLNRNRSIAIPPKMFKLPOQLQHLELRNRIKKNVDGLTFQGLGALKSLKMQR
NGVTKLMGAFWGGLSNMEEILQLDHNNNLTETITKGWLGYLMLQELHLSQNAIRNISPDAWEFC
QKLSLEDDLTTFNHLRSRDDSSFLGLSLLNTLHGNNRVSYIADCAFGRGLSSLKTLDLKNNIEIS
WTIEDMNGAFSGLDKLRRIILQGNRIRSITKKAFTGLDALEHLDLSDNAIMSLQGNAFSQMK
KLOQLHLNTSSLLCDQKLKWLPWVAAENNFOSFVNASCAPHQPLLKGRSIFAVSPDGFWCDDF
PKPQITVQPETQSAIKGSNLSPCSAASSSDSPMTFAWKDNELLHDAEMENYAHLRAQGE
VMEYTTTLLRREVEFASEGKYQCVISNHFGSSYSVKAKLTNVNMLPSFTKTPMDLTIRAGAMA
RLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDFFIVDVKIEDIGVYSCTAQNSA
GSISANATLTVLETPSLPPLDRRTVKGETAVLQCIAAGGSPPPQLNWTKDDSPLVVTERHF
FAAGQNLLIIIVDSDVDSDAGKXTCEMSNTLCTERGVNVRSLVIPTPTCDSPQMTPASLDDDGWA
TVGVVIIAVVCCVVGTSLVWWVIIYHTRRRNEDCSITNTDETNLPADIPSYLSSQGTTLADRG
DGYQVSSESGSHHQFVTSSGAGHFLPQHDSSGTCHIDNSSEADVEATDIFLCPFLGSTGPMY
LKGNVYGSDPFETYHTGCSPDPRTVLMHDHYEPSYIKKCEYPCSHPSEESCRFSNISWPS
HVRKLLNTSYSNHBGPGMKNLCLNKSSLDFSANPEVASVASSNSFMGTFGKALRRPHLDAYS
SFQGPSPDCQPRAFYLKAHSSPDLDSGSEEDGKERTDFQEENHICTFQKQTLNEYRTPNFSYDLDT

Signal sequence:

amino acids 1-27

Transmembrane domain:

amino acids 808-828

N-glycosylation site.

amino acids 122-126, 156-160, 274-278, 442-446, 469-473, 515-519, 688-692, 729-733, 905-909, 987-991, 999-1003, 1016-1020

Glycosaminoglycan attachment site.

amino acids 886-890

Casein kinase II phosphorylation site.

amino acids 99-103, 180-184, 263-267, 314-318, 324-328, 374-378, 383-387, 407-411, 524-528, 608-612, 692-696, 709-713, 731-735, 799-803, 843-847, 863-867, 907-911, 1003-1007, 1018-1022, 1073-1077, 1079-1083, 1081-1085

Tyrosine kinase phosphorylation site.

amino acids 667-675

N-myristoylation site.

amino acids 14-20, 36-42, 239-245, 257-263, 380-386, 427-433, 513-519, 588-594, 672-678, 683-687, 774-780, 933-939

Leucine zipper pattern.

amino acids 58-80, 65-87

FIGURE 107

FIGURE 108

MEGEEAEQPAWFHQWPWRPGASDSAPPAGTMAQSRVLLLLLPPQLHLGPVLAVRAPGFGRS
GGHSLSPPEENEFAEEEPVVLVLSPEEPGPGPAAVSCP RDACSQEGVVDCCGIDLREFPGDLP
EHTNHLSLQNNQLEKIYPEELSRLHRLETLNLQNNRLTSRGLPEKA FEHLTNLNYLYLANNK
LT LAPRFLPNALISVDFAANYLT KIYGLTFGQKPNLRSVYLHNNKLADAGLPDNMFNGSSNV
EV LILSSNFLRHVPKHLPPALYKLHLKNNKLEKIPPGAFSELSSLRELYLQNNYLTDEGLDN
ETFWKLSSLEYLDLSSNNLSRVPA GLPRSLVLLHLEKNAIRSVDANVLTPIRSLEYLLHSN
OLREQGIHPLAFOGLKRLHTVHLYNNALERPVPSGLPRVRVTLMILHNQITGIGREDFATTYF
LEELNLSYNRITS PVQVHRDAFRKLRLRLS LDLSGNRLHTLPPGLPRNVHVLKVKRNEAALA
RGALAGMAQLRELYLTSNRRLRSRALGPRAWVDLAHLQLLDIAGNQLTEIPEGLPESLEYLYL
QN NKISAVPANAFDSTPNLKIGIFLRFNKLAVGSVVDSA FRRLKHLQVLDIEGNLEFGDISKD
RGRLGKEKEEEEEEEEEEETR

Signal sequence:

amino acids 1-48

N-glycosylation site.

amino acids 243-247, 310-314, 328-332, 439-443

Casein kinase II phosphorylation site.

amino acids 68-72, 84-88, 246-250, 292-296, 317-321, 591-595

N-myristoylation site.

amino acids 19-25, 107-113, 213-219, 217-223, 236-242, 335-341,
477-483, 498-502, 539-545, 548-554

Leucine zipper pattern.

amino acids 116-138, 251-273, 258-280, 322-344, 464-486, 471-493,
535-557

FIGURE 109

GGGAGGGGGCTCCGGCGCCGCGCAGCACCTGCTCCGGCGCGCCCTCGCCGTGTCTCCGGGAGGGCGAG
CAGTAGCCCCGGCGGCCACTGGCTGCGCCCAAGGGCTGGGGTTCTCGAGACTCTCACAGGGGGCGCTTCCCACATGGCGCCCAACACCC
CAACCTGTTCTCCCGCTCTACCTGCGGCCCGCCGGAGGTTCAGCGGGAGGTGGCCAGGCAAATAGTCATCGAT
TGCCCTATGCTTATGGTGGGAGGATTGACTGCTGGCTGGGCTGGCTGCCAGTCAGTCAGGCTGGGAGCAGTGTAGCC
TGTGCTGGCAACCCAGATGCAAACATGGTGAATGTCAGGGCCCAAAACAGTGCAGAAGTGTCTACCTGGTTATGCTGG
AAAAACCTGTAAATCAAGATCTAAATGAGTGTGGCTGAAGCCCGCCCTGTAAAGCACAGGTGATAACACTTA
CGGCAGCTACAAGTGTCTACTGTCACCGGATAATGCTCATGCCGGATGGTTCTGCTCAAGTGCCTGACCTG
CTCCATGGCCTAAACTGTCTAGTGGTGGAGGACTGTGTAGATGTGATGAAATGCTGCTCAAGGAAGAGCCTCTGGCCCTAGATT
GCAACTGGCTCTGATGGGAGGACTGTGTAGATGTGATGAAATGCTGCTCAAGGAAGAGCCTCTGGCCCTAGATT
TAGGCACTGTGTCACACATGGGAGGACTCATCTGGCAAGTGTGATGAAATGCTGCTCAAGGAAGAGCCTCTGGCCCTAGATT
CAAATAATATGTCATGCAAGCATGAGCAAACTGGCTACTGGTCAAGTGTGATGAAATGCTGCTCAAGGAAGAGCCTCTGGCCCTAGATT
CGTAGCTGGGCTTACAAGTGAATGAAAGGATACAGGGTGATGGACTGTGACTTGTGTATATTCCTCC
AGITATGTTGAAACCTTCAGGTCCAATCTACATGACCAAGGGAAATGGTACCATTTAAAGGGTGACACAGAAA
TAATAATTTGGATCTCTGATGTTGGAGTACITGGTGGCTCTGACAGAACCATATATTCCTCTATCATTACCA
CAGGCCCTACAGAGCTGAAACACCCTTACCCACCTAACCCAAAGGCCAACACAACTTCCACCAACCCAG
CTGCCCTACAGAGCTGAAACACCCTTACCCACCTAACCCAAAGGCCAACACAACTTCCACCAACCCAG
ACACAGCTGGCAGTACACCTTCAGGGAGGAGGATTACAGTGAACAGGGTACAGACAGACCCCTCAGAAAACCCAGG
AGATGTTGTTGAGTGTGTTGGACACAGTGTGATTTGGATCATGGCTGAGTGGATGGAATCAGGGAGAAAGACAA
TGACTTGACTGGGAACCAATCAGGGACCCAGCAGGTGGACAAATATCTGACAGTGTGCGCAGCCAAAGCCCAGG
GGGAAAAAGCTGAGCTTGGTGTACCTCTGCCGCCCCTGATGCAITCAGGGACCTGTGCTCTGATCAGGCA
CAAGGTGACGGGGCTGCACTTGGCAACTCTGGAGTTGTGAGAAAACACGGTCCACGGAGCAGCCTGTG
GGGAGGAAATGGTGGCCATGGTGGAGGAAACACAGATCACCTTGGAGGGCTGACATCAAAGGCGAATCACA
AAGATGATTAAGGGTTGGAAAAAAAGATCTATGATGTTAAAGGAAACTGGGATTATGGCTTGGAGAAG
AGAAGACTTGAGGGAAACCATGTGTTTCAACTATATGAAGGGTTGGCAAGAGAGGGTGGCAGCACAGCTG
TCTCCATATGCACTAAAGATGAGTGGTTTCAACTATATGAAGGGTTGGCAAGAGAGGGTGGCAGCACAGCTG
GGCCATGTTGAAATCTCTCATAAAAAAAGAAGTGTGAAATCTCATGTTTCAACTATGAGTGGGAGGAAACTT
TAAAGGAAATGGTCTTAAAGATGGTTAAAGGAAACTTACCCAAAGGAAAGTAACAAATTATAGAATTTCCAAA
AGATGTTTGTGATCTACTAGTAGTGTGAAATCTTGAAGACTAAATAATGGCAAGGGTTAAATTAGG
CAITTTCCCTTCTGACCTCTTAATGGAGAGGGATTGAAAGGGGAGGCCACCAAATGCTGAGCTCACTGAAATA
TCTCCCTTATGGCATATAACTGACAGCTGAGATGAAAGGGGAAATAATGAAATTTACTTCTGAGTGGAC
AGATATTTTGTGATCTAGTAAATGCTTGTGAGTGGGTTTCAAGTGTGTTTCAAGTGTGTTCTCATGTAAGGTTAAAGG
TTTCACTTGTGCACTGGATGTTTCAAGTGTGTTTCAAGGAGTCTTCAAGGAAACAGTCAAGAG
ATTTCATCGGGCTCATCTCTCTCTGGTGTGCAAGTGTATCTTGGCTGAGAAAAGAGTGGCTCC
ACACCCGGAGACCTTCTCACCTCATGAGTGTGTTCTTCAAGGAGTCTTCAAGGAAACAGTCAAGAG
AAACAGTAAATTTTGTGAAACATGGTACAATAGGGTACAATAGGCTGTTCTGTCATTTAACCTGGTAAGGGAGGGCTGG
AGGGGGAAAATAATCATTAAGGCTTGTAGTAAACGGCAGAATATAATGGCTGAGTCACTTAAATGGTTCATT
TCTCTTATGGTCACTAAACTGACAGCTGAGATGAAAGGGGAAATAATGAAATTTACTTCTGAGTCCAA
TGATACATGGCTACAATGTGAGGAAAGTATCCAAAGTACTGTATAACATCTTGTGTTTAAATTTAATGGT
CTAAAAAATTTGGTGTGTTGGTTCCAAATGGCTAATAAAACAAATTTGGTAAATAAAACACTGTTAGTAAAT

FIGURE 110

MDFLLALVLVSSLYLQAAAEEFDGRWPRQIVSSIGLCRYGGRIDCCWGWARQSWGQCQPVCQP
RCKHGE CIGPNKCKCHPGYAGKTCNQDLNECGLKPRPCFKRCMNTYGSYKCYCLNGYMLMPD
GSCSSALTCSMANCQYGC DVVKVGQIRCQCPSPGLHLAPDGRTCVDVDECATGRASCPRFRQC
VN TFGSYICKCHKGFDLMYIGGKYQCHDIDECSLGQYQCSSFAR CYNVRG SYKCKCKEGYQG
DGLT CTVYIPKV MIEPSGP IHPVKGNGTILKGDTGN NNWI PDVGSTWWPPKTPVIPPIITNRP
TSKPTTRPTPKPTPIPTPPPPLPTELRTPLPPTPERPTTGLTTIAPAASTPPGGITVDN
RVQTDPQPKPRGDVF SVLVHSCNFHDHGLCGWIREKDNDLHWEP I RD PAGG QYLT VSAAKA PGG
KAARLVLPLGRLMHSGDLCLSPRHKVTGLHSGTLQVFVRKHGAHGAALWGRNGGHGWRQTOI
TLRGADIKSE SQR

Signal sequence:

amino acids 1-17

N-glycosylation site.

amino acids 273-277

Casein kinase II phosphorylation site.

amino acids 166-170, 345-349

Tyrosine kinase phosphorylation site.

amino acids 199-206

N-myristoylation site.

amino acids 109-115, 125-131, 147-153, 191-197, 221-227, 236-242,
421-427, 433-439, 462-468, 476-482

Aspartic acid and asparagine hydroxylation site.

amino acids 104-116, 186-198, 231-243

Cell attachment sequence.

amino acids 382-385

EGF-like domain cysteine pattern signature.

amino acids 75-87

FIGURE 111

CTTCTTTGAAAAGGATTATCACCTGATCAGGGTCTCTGCATTGCCCTTAGTTGATGTGA
AATGTGGCTCAAGGTCTTCACAACTTCCCTTGCACAGGGTCTCGCTGGGGTGA
AGGTGACAGTGGCATCACACACTGTCCATGGCGTCAAGGGTCAAGGCCCTACCTACCGTC
CACTATGGCTTCCACACTCCAGCATCAGACATCCAGATCATGGCTATTGAGAGACCCA
CACAACTGGCCAAATACTTACTGGGCTCTGTGAATAAGTCTGTGGTCTGACTTGGAAATACC
AACACAAGTTACCATGATGCCACCAACTGCACTCGTCTTACACCCACTGCAGTTCCCT
GATGAAGGCAATTACATCGTAAGGTCAACATTCAAGGGAAATGGAACACTATCTGCCAGTCA
GAAGATAACAAGTCACGGTTGATGATCCTGTCAAAAGGCAGTGGTCAAGATTCACTCCCT
CTGGGGCTGTGGAGTATGTGGGAACATGCCCTGACATGTGGAAAGGGGCACTCGG
CTAGCTTACCAATGGCTAAAAAAATGGGACACCTGTCACACAGCTCCACACTACTCTTTC
TCCCCAAAACAATACCCCTCATATTGCTCCAGTAACCAAGGAAGACATTGGAAATTACAGCT
GCCCTGAGGAACCCCTGTCACTGAAATGGGAAATGATATCATTATGCCCATATATATTAT
GGACCTTATGGACTTCAGTGAAATTCTGATAAAAGGGCTAAAGTAGGGGAAGTGTGTTACTGT
TGACCTTGGAGAGGCCATCTTGTGATTGTTCTGCTGATTCTCATCCCCCAACACCTACT
CCTGGATTAGGAGGGACTGACAATACTACATATATCATTAGACATGGGCTTAGAAGTT
GCATCTGAGAAAATGGCCAGAACAGACATGGACTATGTGCTGTGCTTACAACAAACATAAC
CGGAGGCAAGATGAAACTCATTCACTGAGTATCATCCTCGTAGGACTGAGAACGCTTG
CACAGAAAGGAAAATCATTGTCACTTCTAGCAAGTATAACTGGAATATCACTATTGGT
ATATCCATGTGCTTCTCTTCTGAAATGGAAAATTCAGACCTACAAAGTTATAAAACAGAA
ACTAGAAGGGCAGGCCAGAAAACAGAAATCAGGAAAGCTCAAACATTTCAGGCCATGAAGATG
CTCTGGATGACTCGGAATATATGAATTGTTGCTTTCAGATGTTCTGGTGTTCAGG
ATTCCAAGCGAGGTCTGTTCCACCCCTGATGTGATCAGGGGAAGATTGCAAGACTACAGT
GTGAGGTTTACAGCACATCCCTGCCAGCAGAACGACATCCAGAGTGAACCTTCATGG
GCTAACAGTACATTGAGTGAAAATTCTGAGAAAACATTAAAGGGAAAACAGTGAGGAAAGT
ATATTAATCTGGAATCAGTGAAGAAACCAGGCCAACACCTTACTCATTATCCCTTACA
TGCAGAAATAGAGGCCATTATGCAAAATTGAACTGCAGGTTTTCAGCATATACACAATGTCTT
TGCAACAGAAAACATGTTGGGAAATTCTCAGTGGAGAGTCGTTCTCATGTCAGGG
GGAGAACGAAAGTGACAGGGTTCTCATTAAGTTTGATGAAATATCTCACAAACCTCA
ATTAGTTCTACTCTACACTTCACTTCAACACTGAGACTATCCTGTCACCTACAAA
TGTGGAAACCTTACATTGTTGATTTTCTCAGCAGACTTTGTTTATTAAATTTTTATTAGTG
TTAAGAATGCTAAATTCTACATTGTTCAATTTATTCAGCAACTTGTGTTTATTGACAA
CAAAGTAATAAGGATGGTTGTCACAAAACAAACTATGCCCTCTCTTTTTTCAATCACC
AGTAGTATTGGAGAAGACTTGTGAACACTTAAGGAAATGACTATTAAAGTCTTATTGTTA
TTTTTCAAGGAAAGATGGATTCAAAATAATTCTGTTTGTCTTTAAAAAAA

FIGURE 112

MWLKVFTTFLSFATGACSGLKVTVPSPHTVHGVRGQALYLPVHYGFHTPASDIQIIWLFERPH
TMPKYLLGSVNKSVVVPDLEYQHKFTMMPPNASLLINPLQFPDEGNYIVKVNIQGNGLSASQ
KIQVTVDVPVKPVVQIHPPSGAVEYVGNTLTCHEGGTRLAYQWLKNGRPVHSTSSTYSFS
PQNNTLHIAPVTKEDIGNYSCLVRNPVSEMESDIIMPIIYYGPYGLQVNNSDKGLKVGEVFTV
DLGEAIALFDCSADSHPPNTYSWIRRTDNNTYIIKHGPRLEVASEKVAQKTMVDYVCCAYNNIT
GRQDETHEFTVIIITSVGLEKLAQKGKSLSPASITGISLFLIISMCLLFLWKKYQPYKVIKQK
LEGRPTEYRKAQTFSGHEDALDDFGIYEFVAFPDVSGVSRIPRSRVPAASDCVSGQDLHSTV
YEVIQHIPAQQQDHPE

Signal sequence:

amino acids 1-18

Transmembrane domain:

amino acids 341-359

N-glycosylation site.

amino acids 73-77, 92-96, 117-121, 153-157, 189-193, 204-208,
276-280, 308-312

Casein kinase II phosphorylation site.

amino acids 129-133, 198-202, 214-218, 388-392, 426-430, 433-437

Tyrosine kinase phosphorylation site.

amino acids 272-280

N-myristoylation site.

amino acids 15-21, 19-25, 118-124, 163-167, 203-209, 231-237,
239-245

Prokaryotic membrane lipoprotein lipid attachment site.

amino acids 7-18

FIGURE 113

GCAAGCGGCCGAAATGGCGCCCTCCGGGAGTCTTGCAGTTCCTGGCAGTCTGGTGTGTT
GCTTGGGGTGTCCCCTGGACGACGGCGGGAGCACGTTCGCTCATCAGGACGAGA
ACTGGAGAGAAACTGCTGGAAGGAGACTGGATGATAAGAATTTCATGCCCTGGTGTGCTC
TGTC_{AA}ATCTCAACCGGAATGGGAAAGTTTGCTGAATGGGAGAAGATCTTGAGGTTAA
TATTGCGAAAGTAGATGTCACAGAGCAGCCAGGACTGAGTGGACGGTTATCATAACTGCTC
TTCCCTACTATTATCATTGTAAGATGGTGAATTAGCGCTATCAGGGTCAAAGGACTAAAG
AAGGACTTCATAAACTTATAAGTGATAAAAGGTGGAAGAGTATTGAGGCCGTTCATCATG
GTTTGGTCCAGGTTCTGTTCTGAGTGAGTACTATGTCAGCACTCTTCAGCTATCTATGTGGA
TCAGGACGTGCCATAACTTATTGAAGACCTTGGATGCCAGTGGGGATCATATACT
GTTTTCCTTAACTGTCAGGACTGTTTCCGGACTGTTATTAGGACTCTGTATGATAATTGTC
AGATTGCCCTTGCTTCAAAAAGCGCAGACCACAGCCATACCCATACCCCTTCAAAAAAAT
TATTATCAGAAATCTGCAACCTTGTAAAAAGTGGAGGAGAACAGAGGGGATGAAAGAA
GATGTTTCAGAAGAAGCTGAAAGTAGTAAAGAAGGAAACAACAAAGACTTCCACAGAATGC
CATAAGACAAACGCTCTGGGCCATCATTGGCCACAGATAATCTTAGTTAAATTTATAG
TTATCATAATTATGATTGTAAAAAAGAGATGATCATTGTTGGTTGAGTG
AACTGTGACTTTTGTAAATTGCGGGTTCTGACTGATGTTGATTAATTGAGACTCTA
CATTCAAGACATAAAAGCACTTAGTTGATACAAGTTGAAATATTGATTAAGCACAGTATGATG
GTTAAATAGTCTCTAATTGAAAAATCTGTCAGGTTGATTAATTGATTAATTGATTAATTGAT
TTAATAAACTTCAATTGAGCTGAGTGTGAAATTTGACATTTCCAAGTATTGCAATTAT
TGAGGTTATTAAAGAAAGATTATTGAGAAAAATATTCTCATTTGATATAATTCTCTG
TTCACTGTGGGAAAAAGAGATAATTCCCATATAATTGGAAGTTGCCATTGTC
AAATGTTGATTTCAGTGACAATTCTGGTCTTTAGAGGTATATTCCAAAATTCTCTG
ATTTTAGGTATTGCAACTATAAAACACTTACCTTACATTAAATTACAGTTTCTACACA
TGTTAATACAGGATATGCTACTGATTAGGAAAGTTTAAGTTCATGGTATTCTCTGATT
CAACAAAGTTGATTCTCTGTATTGTTCTACTATGTTGATTAACAGTATTGTTGTT
CAAATTGGATGATAATTGCGAAACATTTTGATTTGTTGATTAACAGTATTGTTGTT
GTTTCAAAACTGAAGTTACTGAGAGATCATCAAAATTGACAACATTGTTGTAATTAA
AAATTTTGGCCACTTTTCAGATTACATCATTCTGCTGAACCTCAACTGAAATTGTTTTT
TTTCTTTGGATGAGGTGAAGGTGAACATTCTGATTTTCTGATTTCTGATGTTGAAAAAGCCTGGTA
TTTACATTGAAATTCAAGAGCTTAATATAAAAGTTGCTCTCATATCAGAAAGTCTTAATTGAT
TTACAGTCTGTAATGCTTGATGTTTAAATAAAACATTTTATATTTTAAAGACAA
ACTTCATATTATCTGTTCTTCTGACTGGTAATATTGTTGCGGATTTCACAGGAAAAA
GTCAGTAGGATGGAAACATTTACTGTAATTCTACTCTTAAGAGCTAGAATACATAGTTT
CACCTTAAAGAAGGGGAAATCATAAATAACATGAATCAACTGACCAATTACGTAGTAGAC
AATTTTCTGTAATGCCCCCTTCTAGGCTCTGTTGCTGTGAATCATTAGATTACAG
TATCGTAATATAACAGTTCTTTAAAGCCCTCTCTTCTAGAATTAAATATTGACCTT
AAAGAGTTGGATGCTAACTGCTGATGCCCTAGAAAAATATCTAACGACAAAATAAACCT
TTCTAACCACTTCATTAAAGCTGAAAAAAAAAAAAAA

FIGURE 114

MAPSGSLAVPLAVLVLLWGAPWTHGRRSNVRVITDENWRELLEGDWMIIFYAPWCPACQNL
QPEWESFAEWGEDLEVNIAKVDVTEQPGLSGRFIITALPTIYHCKDGEFRRYQGPRTKKDFI
NFISDKEWKSIEPVSSWFGPGSVLMSSMSALFQLSMWIRTCHNYFIELDLGLPVWSYTVFAL
ATLFSGLLLGLCMIFVADCLCPSKRRRPQPYPPSKLLSESAQPLKKVEEEQEADEEDVSE
EEAESKEGTNPKDFPQNAIRQRSLGPSLATDKS

Signal sequence:

amino acids 1-26

Transmembrane domain:

amino acids 182-201

Casein kinase II phosphorylation site.

amino acids 68-72, 119-123, 128-132, 247-251, 257-261

Tyrosine kinase phosphorylation site.

amino acids 107-115

N-myristoylation site.

amino acids 20-26, 192-198

Amidation site.

amino acids 25-29

FIGURE 115

GCAGGTGTCAGCTCGGGAGACCCGTGATAATTCTGTTAACTAATTCAACAAACGGGCCCTT
CTGTGTGCCAGAACCGCAACGAGCTTGCTAACCCAGTGGACAGGCGGATTGGAAGAGCGGG
AAGGTCTGCCAGAGCAGTGTGACACTTCCCTCTGACCATGAAACTCTGGGTGTC
ATTGCTGATGCCCTGGTTGGTGTCTGAGCTGTGTCAGGCGAATTCTTACACCTCTATTG
GGCACATGACTGACCTGATTATGAGAGAAGAGCTGGTGCAGTCTCTGAAAGAGTACATC
CTTGTGGAGGAAGCCAAGCTTCAAGATTAAAGAGCTGGCCAACAAAATGGAAGGCTTGC
TAGCAAGTCAGCTGCTGATGCTGAGGGCTACCTGGCTCACCTGTGAATGCCCTACAAACTGG
TGAAGCGCTAAACACAGACTGGCTGCGCTGGAGGACCTTGTCTGCAGGACTCAGCTGCA
GGTTTATCGCACCCTCTCTGTCAGGCGAGTCTTCCCCACTGTAGGGACGAGATAGG
AGCTGCCAAAGCTGATGAGACTTCAGGACACATAAGGGCTGGACCCAGGCAATTCTCA
GAGGGGAACCTCCAGAACCAAGTACCAAGGAATGCTGAGTGTGGATGACTGCTTGGGATG
GGCGCTCGGCCAACATGAGAGGACTATTATCATCGCTGTGGATGGAGCAGGTGCT
AAAGCAGCTGTGATGGGACTACAGGACCAACAAAGCTCACAGGTGCTGACTACCTCA
GCTATGCTGCTTCAGTTGGGTGATCTGCACCGTGCCCTGGAGCTCACCGCCGCTGCTC
TCCCTTGACCAAGCCAGCACAGAGCTGGAGGAATCTGGGTACTTTGAGCAGTATTGGA
GGAAAGAGAGAAAAAACGTTAACAAATCAGACAGAACGCTGAGCTAGCAACCCCAGAAGCA
TCTATGAGAGGCCCTGTGACTACTCTGCTGAGGGATTTACAGAGAGCCTCTGCTGGG
GAGGGGTCAAACACTGACACCCCGTAGACAGAAAGAGGCTTCTGTAGGTACCACTGGCAA
CAGGGGCCAACAGCTGCTATGGCCCTTCAAAGAGGGAGGAGCTGGGACAGCCCGCACA
TCTGAGGTACTACGATGTCATGTCAGGAAATCGAGGAGATCAAGGAGATCGCAAAA
CTCAAACCTGCAAGGCCACCGTGTGATCTCAAGACAGGAGTCTCACTGTCGCCAGCTA
CCGGGTTTCCAAAAGCTCTGGTAGAGGAAGAGTGTGACCTGTTGGCCCGAGTAAATC
GTGGGATGCGACATCACAGGGTTAACAGTAAAGACTGAGAATTGTTACAGGTTGCAAAT
TATGGGACTGGGAGGACAGTATGAAACCGCACTTCGACTTCTGAGGGACCTTTGACAGCGG
CCTCAAAACAGAGGGAAATAGGTAGCGACGTTCTTAACTACATGAGTGTAGAAGCTG
GTGGGTGCAACCGTCTCCCTGATCTGGGGGCTGCAATTGGCCAAGAAGGGTACAGCTGT
TTCTGGTCAACCTCTGGGAGCCGGGAAGGTGACTACCGAAAGACATGTCGCCCTGCC
TGTGCTTGTGGCTCAAGTGGGTCTCCAAATAAGTGGTCCATGAACGAGGACAGGAGTTCT
TGAGACCTTGTGGATCAACAGAAGTTGACTGACATCCTTTCTGCTCTCCCTTCCCTGGTC
CTTCAGCCCATGTCACAGTGCAGACAGACACCTTGTGATGTTCTGATGTTCTATCAGGCT
GATTTTGGAGAAATGAATGTTCTGTGGAGCAGAGGGAGACCATACTAGGGGACTCTGT
GTGACTGAAGTCCCAGCCCTTCATTCAAGGCTGTGCCATCCCTGGCCCAAGGCTAGGATCA
AAAGTGGCTGCAGAGTTAGCTGTCTAGCGCCTAGCAAGGIGCCTTGTACCTCAGGTGTT
TTAGGTGTGAGATGTTCTAGTGAACCAAGTTCTGATACCTTGTGTTACATGTTGTTTAT
GGCATTTCTATCTATTGTGGCTTACCAAAAATAAAATGTCCCTACAGAAAAAA

FIGURE 116

MKLVVSALLMAFGVLSCVQAEFFTSIGHMTDLIYAEKELVQSLKEYILVEEAKLSKIKSWA
NKMEALTSKSAADAEGYLAHPVNAYKLVKRLNTDWPALEDLVLQDSAAGFIANLSVQRQFFF
TDEDEIGAAKALMRLQDTYRLLPGTISRGEGLPGTKYQAMLSVDDCFGMGRSAYNEGDDYHTV
LWMEQVLKQLDAGEEATTTKSQVLDYLSYAVFQLGDLHRALELTTRRLSLDPSHERAGGNLR
YFEQLLEEEREKTLTNQTEAELATPEGIYERPVVDYLPERDVYESLCRGEGVKLTPRRKRLF
CRYHHGNRAPQLLIAPFKEEDEWDSPHIVRYYDVMSDEEIERIKEIAKPKLARATVRDPKTG
VLTVASYRVSKSSWLEEDDDPVVARVNRMQHITGLTVKAELLQVANYVGCGQYEPHDFDS
RRPFDSGLKTEGNRLATFLNYMSDVEAGGATVFPDLGAAIWPKKGTAVFWYNLLRSGEQGDYR
TRHAACPVLVGCKWSNWKWFHERGQEFLRPCGSTEVD

Signal sequence:

amino acids 1-17

N-glycosylation site.

amino acids 115-119, 264-268

Glycosaminoglycan attachment site.

amino acids 490-494

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 477-481

Casein kinase II phosphorylation site.

amino acids 43-47, 72-76, 125-129, 151-155, 165-169, 266-270,
346-350, 365-369, 385-389, 457-461, 530-534

Tyrosine kinase phosphorylation site.

amino acids 71-80, 489-496

N-myristoylation site.

amino acids 14-20, 131-137, 171-177, 446-452

Prokaryotic membrane lipoprotein lipid attachment site.

amino acids 8-19

Leucine zipper pattern.

amino acids 213-235

FIGURE 117

GCAGTATTGAGTTTACTCTCCTCTTTAGTGGAAAGACAGACCAATACTCCAGTGAGTGAATTGATGT
TTCATTTAACCGTTTGGCTGGGGTAGTCCGACACCTTCAGATGAAGAGCAGGCCAGAGAGATGTGAGA
AGACAGGACAATCTCTGGGGATGCTGGCTCTGGAAAGCCAGGGGCTTGCTCTGCTCTGCTCTGGCTCAT
CAGGTTCTCTGGTAAAGCTACTACTGCGCTGGGCTCATTAATCCATTGATCCTTGAGGCTGTGCC
CTCTGGCTCACCCACCTGAGGCCAGGGCTTACACCATGAGCTGAGCTCCCTGTTGCTCTGCGCCAGGCC
CCCTCATCTTGGCTGTCTGGGCTGAGGCCAGGGCTTACAGGGACCCCACAAGGCCATACAGAAGGTGCT
ATGCCAGGCTTACAGGGCTTACAGGGACCCCACAAGGCCATACAGAAGGTGCTCAGGCC
ATGAGAGACTTCAAACCCGGATTGTCCTACTACAGGGACCCCACAAGGCCATACAGAAGGTGCT
GGTACATCCAGACAGAGCTGGGCTCCGGTGGAGCTGCTGGTGGGCTGCTGACCTCCCGAGCTACATGTC
CTTGGGCCGTGCGCTGTGACACCGTAGCGCTGGGCTACATTCCTCGTACTACTCTACTCTACT
CCCCGGGCTTCCAGCAGGGATGCAGGGTGTCTCATGGGATGAGCGCCGGGCTGTGCTATGTCAGAGACCTTC
GCCACCTTCACACACTTGGGGCCGACTACAGACTTCTCTGGGCCACTCAGCATCAACAAAGACCTGTACT
CCCCCTGGCAGCCCTGCTGGGCCACTCAGCATCAACAAAGACCTGTACTTAGGCCGGCAGAGGATCTATTG
GCCAGGCGAGACGGCCGGGACTCTGATGGGGGATGAGCGCCGGGCTGTGCTATGTCAGAGACCTTC
GCCACATCTGAGATGGCTGGAGGAGACATCTCAGTGGCTCCCTCTGAGCAGTGGCTTGGAGCTGCTCAT
ACTCTCTGGGGCTCGGGTGTCTCACAGCACAAAGGGCAGCAGTACTCTGCTCATTTGAACTGGCCAAA
ACCTGAGAAAGGAAAGGAGACTGGGCTTCTCTGGGCTTCTCTGGGCTTCTCTGGGCTTCTCTGGGCT
ACCGGCTCCACAAACCTCTCACGGTCTGGGAGTGGAGCCGGCTTAACTGAAATAGAACACTGCA
TCCGAGGACTGACCGTGTGACCCGGAGGG
CACCAACACTCTCGGCTTGGGGCTGCTGGGACTACTCTCACAGAGCAGCACACCTTCTCTGAGCAGATGGG
CTCCCCAAGTGGGACTACAGGGGGTGTGAGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG
ATCGTCAGGCTATCACGGCCGGCTGGCTCCCTGGGAGGAGCTGCTGGGTGAGCTGGGAGACTG
GGGGCATGGAGTAACCTGGACACTGGCTCTGGGAGGAGCTGCTGGGAGGAGCTGGGGGGGGGGGGGGGG
GGGTGAGGGCTGG
AGCTGG
TGGAGCCAGAGAACATGCAATTGCTACCTGGCTTACGGGGCACAGAGAAAGTGGCCGTGGAGCTCCAG
ACCCATTCTGGGGTAGGGCTGAGCAGCGGAGTTAGAGGAGCAGCTGGGGGGGGGGGGGGGGGGGGGGGG
CTGG
TCTTCTTCTTACACCGTGTGGAACAGGCTGG
GGCAGGGCTTCTTCTGG
GG
ACCCGGCAGGGCTTCTGG
TGGCAGGGCCAGGAAGAGGGAGGAAGGGCCCTGGAGGGGGCTGGAGGGGGTGTGGATGTTTCTCCGG
ACCTCTTCTGG
ACCTTACCAACCGCTGGCCGCTCACACCTGGAGGGGGTAGGGGGGGGGGGGGGGGGGGGGGGGGGGGG
AGCAGGGAGCAGGCCAATAGCACTTAGCCGGCTGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG
CCAGGAAGGGCAAGGCAAGGCAAGGAGATGGGAGAGATAGAGAAATTGTTGCTGTTATTTTAAAT
ACATGTCCTCTGCC

FIGURE 118

MRLSSLLALLRPALPLI LGLS LGCSLSLLRV SWIQGE GEEDPC VEA VGERGGPQNPDSRARLD
QSDEDFKPRIVPYRDPNKP YKKV LRT RYI QTELGS RER LLV A VLTS R ATLST LAV AVNRTV
AHHF PRLLYFTG QRGARAPAGM QVSHG DERPAWLMSETL RHLH THFGADYDWFFIM QD DTY
VQAPRLA ALAGHLSIN QDLYL G RAE F I GAGE QARYCHGGFGYLLSRSLLLRLP HLDGCRG
DILSAR PDEWLGRCLIDSLGVGC VSQHQGQYR SFELAKNRDPEKEGSSAFLSAFAVHPVSE
GTLMLYR LHKRFSALELERAY SEIEQLQ A QIRNL TVLTP EAGL SWFV GLPAPFTPHSRF EV
LGWDYFT EQHTFSCADGAPKCP LQGASRADVGDALETALEQLNRRYQPRLRFQ KQRLLNGYR
RFPD PARGMEY TLDL LECVTQ RG HRRALARRVSLLRPLSRVE ILPMPYVTEATRVQLVLPLL
VAEAAA APAFLEA FAANVLEPREH ALLTLLVYGPREGGRGAPDPFLGVKA AAAE LERRY PG
TRLAWLAVRAEAPS QVRLMDVVSKKH PVDTLFLTTVWTRPGPEV LNCRMNAI S G WQ AFFF P
VHFQE FNPA LSPQRSPGPPGAGPDPPSPGADPSRGAFIGG RFD RQAS AEGCF YNADYI AA
RAR LAGELAGQ EEEA LEGLE VMDVFLRF SGLHL FRAVE PGLVQKF SLRDCSP RLSE ELY HR
CRLSNLE GLG GR AQLAMALF E QEQ ANST

Signal sequence:

amino acids 1-15

Transmembrane domain:

amino acids 489-507

N-glycosylation site.

amino acids 121-125, 342-346

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 319-323, 464-468

Casein kinase II phosphorylation site.

amino acids 64-68, 150-154, 322-326, 331-337, 368-372, 385-389,
399-403, 409-413, 473-477, 729-733, 748-752

Tyrosine kinase phosphorylation site.

amino acids 736-743

N-myristoylation site.

amino acids 19-25, 23-29, 136-142, 397-403, 441-447, 544-550,
558-564, 651-657, 657-663, 672-678

Prokaryotic membrane lipoprotein lipid attachment site.

amino acids 14-25

Cell attachment sequence.

amino acids 247-250

FIGURE 119

CGGAGTGGTGCGCCAACGTGAGAGGAACCGTGCGCGCTGCGCTTCCTGCCCCAAGCC
GTTCTAGACGGGGAAAAATGCTTCTGAAAGCAGCTCCTTTGAAGGGTGTATGCTTGG
AAGCATTTCGTGCTTGATCACTATGCTAGGACACATTAGGATTGGTCAATGAAATAGAA
TGCACCACCATGAGCATCATCACCTACAAGCTCTAACAAAAGAAGATATCTTGAAGATTTCA
GAGGATGAGCGCATGGAGCTCAGTAAGAGCTTCGAGTATACTGTATTATCCITGTAAAACC
CAAAGATGTGAGTCTTGGGCTGAGTAAGGAGACTGGACCAAACTGTGACAAAGCAG
AGTTCTTCAGTCTGAAAATGTTAAAGTGTGAGTCATTAAATATGGACACAAATGACATG
TGGTTAATGATGAGAAAAGCTTACAAAATACGCTTGTATAAGTATAGAGACCAATACAACTG
GTTCTTCCTTGCACGCCACTACGTTGTATCATTGAAAACCTAAAGTATTTTGTAA
AAAAGGATCCATCACAGCCTTCTATCTAGGCCACACTATAAAATCTGGAGACCTGAAATAT
GTGGGTATGGAAGGAGGAATTGCTTAAGTGTAGAATCAATGAAAAGACTTAACAGCCTTCT
CAATATCCCAGAAAAGTGTCTGAAACAGGGAGGGATATTGGAAGATATCTGAAGATAAAC
AGCTAGCAGTTGCCTGAAATATGCTGGAGTATTGAGAAAATGCAAGAGATGCTGATGGA
AAAGATGTATTAATACCAAATCTGGGGCTTCTATTAAAGAGGCAATGACTTATCACCC
CAACCAGGTAGTAGAAGGCTTGTTCAGATATGGCTGTTACTTTAAATGGACTGACTCCAA
ATCAGATGCATGTGATGATGTATGGGTATACCGCTTGGCATTGGGATATTTCATTTCAAT
GATGCATTGGTTCTTACCTCCAATGGCTGACAATGACTTGAGAAGTGGTAGAAAAGCG
TGAATATGATCTTGTATAGGACGTGTGTGTCATTGGTAGTAACTACATATCCAA
TACAGCTGTATGTTCTTTCTTCTAAATTGGTGGCACTGGTATAACCACACATTAAG
TCAGTAGTACATTTAAATGAGGGTGGTTTTCTTAAACACATGAACATTGAAATG
TGTGGAAAGAAGTGTGTTAAGAATAATAATTGCAAAATAACTTAAATAATATTATAT
GTGATAAAATTCTAAATTATGAAACATTAGAAATCTGTGGGCACATATTGCTGATGGTT
AAAAAAATTAAACAGGTCTTAGCGTTCTAAGATATGCAAATGATATCTCTAGTTGTGAATT
TGTGATTAAAGTAAAACCTTTAGCTGTGTGTTCCCTTACTCTAAACTGATTATGTTCT
AAGCCTCCCCAAGTCCAATGGATTGCTTCAAAATGTACAACTAAGCAACTAAAGAAA
ATTAAGTGAAGTGTGAAAAT

FIGURE 120

MLSESSSFLKGVMLGSIFCALITMLGHIRIGHGNRMHHHEHHHLQAPNKEDILKISEDERME
LSKSFRVYCIILVKPKDVSLWAAVKETWTKHCDKAEFFSSENVKVFESINMDTNDMWLMMRK
AYKYAFDKYRDQYNWFFLARPTTFAIIENLKYFLLKKDPSQPFYLGHТИKSGDLEYVGMEGG
IVLSVESMKRLNSLLNIPEKCPEQGGMIWKISEDKQLAVCLKYAGVFAENAEDADGKDVFNT
KSVGLSIKEAMTYHPNQVVEGCCSDMAVTFNGLTPNQMHVMMYGVYRLRAFGHIFNDALVFL
PPNGSDND

Signal sequence:

amino acids 1-33

N-glycosylation site.

amino acids 121-125, 342-346

cAMP- and cGMP-dependent protein kinase phosphorylation site.

amino acids 319-323, 464-468

Casein kinase II phosphorylation site.

amino acids 64-132, 150-154, 322-326, 331-335, 368-372, 385-389,
399-403, 409-413, 473-477, 729-733, 748-752

Tyrosine kinase phosphorylation site.

amino acids 736-743

N-myristoylation site.

amino acids 19-25, 23-29, 136-142, 397-403, 441-447, 544-550,
558-564, 651-657, 657-663, 672-672

Prokaryotic membrane lipoprotein lipid attachment site.

amino acids 14-25

Cell attachment sequence.

amino acids 247-250

FIGURE 121

CCACAGCGTCCGATCTTACCAACAAAACACTCCTGAGGGAGAAAGAGAGAGGGAGGGAGAG
AAAAAGAGAGAGAGAGAAACAAAAACCAAAGAGAGAGAGAAAAATGAATTCTAATCAT
CTGAAACACAATGCACAGAGAGAGGATGCTCTTCCCAAATGTTCTATGGACTGTTGCT
GGGATCCCCATCCCTATTCAGTGCTGTTCATCACAGATGTGTTGACATTTCGCAT
CTTCAAAACCTGTGATGAGAAAAGTTCAAGTACACTGAGAATTTCACAGAGCTCTGCT
ACAATTATGGATCAGGTTCAAGTCAAGAATTGTTGTCATTGAACTGGGAATATTTCAATCC
AGCTGCTACTCTTTCTACTGACACCATTCTGGCGTTAACGAGGGAGCAGGAATTCTTCAAGA
CATGGGGCTCACCTGGGGTATCAACTCACAGGAGGAGCAGGAATTCTTCAAGA
AACCTAAAATGAGAGAGTTTTATTGGACTGTCAGACAGGTTGTCAGGGTCAGGGAA
TGGGTGGACGGCACACCTTGACAAAGTCTCTGAGCTTCTGGATGTAGGGAGCCAAACAA
CATAGCTACCCCTGGAGGACTGTGCCACCATGAGAGACTCTCAAACCCAAAGGCAAATTGGA
ATGATGTAACCTGTTCTCAATTATTTGGGATTTGTGAAATGGTAGGAATAATCCTTG
AACAAAGGAAATCTTTTAGAACAGAACGGCACAACCTCAAATGTTAAAGAAGGAAGAGCA
AGAACATGGCACACCCACCGCCCCACAGAGAAATTGCGCTGAACCTCAAAGGACTTC
ATAAGTATTGTTACTCTGATACAAATAAAATAAGTAGTTAAATGTTAAAAAAAAAAAA
AA
AAAAA

FIGURE 122

MNSSKSSETQCTERGCFSSQMFLWTVAGIPILFLSACFITRCVVTFRIFQTCDEKKFQLPEN
FTELSCNYGSGSVKNCCPLNWEYFQSSCYFFSTDТИWALSLKNCSAMGAHLVVINSQEEQ
EFLSYKKPKMREFFIGLSDQVVEGQWQWVDGTPLTKSLSFWDVGEPNNIATLEDCATMRDSS
NPRQNWNNDVTCFLNYFRICEMVGINPLNKGKSL

Signal sequence:

amino acids 1-42

N-glycosylation site.

amino acids 2-6, 62-66, 107-111

Casein kinase II phosphorylation site.

amino acids 51-55, 120-124, 163-167, 175-179, 181-185

N-myristoylation site.

amino acids 15-21, 74-80, 155-161

Prokaryotic membrane lipoprotein lipid attachment site.

amino acids 27-38

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FIGURE 123

GGGATAACAGCGCCGCCTGGCCCCCTCAGCACCCCTCGACATGGCGCTGAGGGGCCACCGCGAC
TCGGGCTCTCGCTCGCTGACTCTCTCTCTGCTCTTCAAGGGCTGCCATAGGGCTGTAAATC
TCAAATCCAGCAATCGAACCCCCAGTGGTAACAGGAATTGAAAGTGGAACTGTCTGCATCATTAAGGATTGCG
AGACAAGTCGGGCAAGGGAGACTGGTAACAGGAATTTCAGAGTGAACACACATATGTTTTTGACAAACAAA
TTCAAGGGAGACTTGGCGGGCTGTCAGAAAATCTGGGAGAGACATCCCTGAAGATCTGGAAATGTGACACGGAG
ACTTCAGGGCCTTATTCGCTGTGAGGTCGTTGCTGAGGAAATTGATGAGATTGTGATCGAGTTAA
CTGTGCAAGTGAAGCCAGTGACCCCCCTGCTGAGAGTGCAGGCTGTACCGAGTAGGCAAGATGCCAACACTGC
ACTTCAGGGAGACTGGGGCCACCCCGGCCCTACATACAGCTGGTATGCCAATGTGATGACCCACTGCCAACCGGATT
CCAGAGCCAACTCCAGATTTCGCAATTCTCTTCCACTAAACCTGAAACAGGCACCTTGGTGTTCATGCTG
TTCAAAAGGACGACTCTGGCACTACTGCACTGCTTCAATGACCCAGGCTCAGGAGGTGTGAGGAGCAGG
AGATGAGGAACTTGGGCAATCTGGGAATTGGGGGGTTCTGGTGTCTGCTGTACTGGCCCTGA
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